Medical Officer's Review of NDA 20-803

Original

NDA #20-803 Submission: 1/31/97
M.O. Review #1 Review completed: 8/11/97
Revised printing: 1/27/98

Proposed Tradenames:

Alrex or Altrin

Generic name:

Loteprednol etabonate ophthalmic suspension, 0.2%

Chemical name:

Chloromethyl-17\alpha-[(ethoxycarbonyl-oxy]-11\beta-hydroxy-3-

oxoandrosta-1,4-diene-17 carboxylate

Sponsor:

Pharmos Corporation

2 Innovation Drive, Suite A

Alachua, FL 32615

Pharmacologic Category:

Steroid

Proposed Indication(s):

Treatment of signs and symptoms of seasonal allergic

conjunctivitis

Dosage Form and

Route of Administration:

Ophthalmic suspension for topical ocular administration

NDA Drug Classification: 1S

Related INDs:

Related NDAs:

NDA 20-583 Loteprednol etabonate ophthalmic suspension, 0.5%

NDA 20-841 Loteprednol etabonate ophthalmic suspension, 0.5%

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3 Material Reviewed

NDA 20-803 Volumes 1.1, 1.16-35 NDA 20-583 Studies by reference - See Medical Officer's Review (MOR)

4 Chemistry/Manufacturing Controls - see Chemistry Review

Raw Material	Quantity mg/mL	% label excess	Range
Loteprednol etabonate			
Povidone USP			
Benzalkonium Chloride.		_	
Edetate disodium			
Glycerin			
Tyloxapol			
Purified water	QS to 1 mL		
Sodium Hydroxide	Adjust pH		
Hydrochloric acid '	Adjust pH _		

Additional Specifications:

pН

Osmolality

250-310

Particle size

Sterility

USP

Preservative efficacy

USP

Reviewer's Comments:

- I. Issues related to water loss and the formation of "aggregate" material after storage of inverted containers will need to be resolved prior to approval.
- . 2. The pH range in the NDA summary differs from other sections of the NDA. The range should be clarified.

5 Animal Pharmacology/Toxicology - See Pharmacologist's Review No additional issues identified.

6 Clinical Background

See MOR of NDA 20-583

6.1 Relevant human experience

No previous human experience.

6.3 Foreign experience

No foreign marketing experience. No pending

foreign applications.

6.4 Human Pharmacology

Pharmacokinetics & pharmacodynamics:

See MOR NDA 20-583

7 Description of Clinical Data Sources

Review Number	Protocol	Indication	Design	Treatment Arms	Number in each arm	Age Range	% (d/8) B/W/O	Duration of treatment
1	143	Allergic Conjunctivitis	Parallel Double masked	Loteprodnol Vehicle	66 67	23-73	(50/50) 0/64/39	42 days
2	144	Allergic Conjunctivitis	Peraliel Double masked	Leteprednol Vehicle	67 68	19-74	(46/54) 0/67/32	42 days
3	141	Allergic Conjunctivitis	Paired eye Double masked	Loteprednol	60	19-85	(55/45) 0/97/3	28 days
4	145	Allergic Conjunctivitis	Paired eye Double masked	Lotoprod 0.1% Lotoprod 0.2% Lotoprod 0.3% Lotoprod 0.5%	28 31 29	19-66	(51/49) 0/99/1	28 days

8 Clinical Studies

8.1 Indication # 1 Seasonal Allergic Conjunctivitis

8.1.1 Study #1 Protocol # 143

Title: Safety and efficacy of loteprednol etabonate in the treatment of

seasonal allergic conjunctivitis (QID dosing).

Objective: To evaluate the efficacy and safety of loteprednol etabonate 0.2%

ophthalmic suspension in the treatment of signs and symptoms of

environmental seasonal allergic conjunctivitis.

Study Design: A randomized, double-masked, placebo controlled, parallel

group multicenter (3 sites) study.

Test Drug Schedule: All subjects received either loteprednol etabonate 0.2%

ophthalmic suspension (LE), or placebo (vehicle) bilaterally,

QID for 42 days.

Investigators: Number of Patients Enrolled:

Steven J. Dell, M.D.(#174) 34

Eye Care Austin 1700 S. Mo-Pac Austin, TX 78746

George M. Lowry, M.D.(#175) 60

Vision Care 8123 Broadway

San Antonio, TX 78209

James A. Northcutt, M.D. (#178)

Northcutt Eyecare Center 903 South W.W. White Rd. San Antonio, TX 78220

Study Plan

This was a prospective, double masked, placebo controlled, multi center (3), study in patients with signs and symptoms of seasonal allergic conjunctivitis (SAC). Enrolled in the study were one hundred and thirty three (133) subjects, with a history of positive skin prick or RAST test, and at the time of enrollment, presenting with moderate to severe signs and symptoms of seasonal allergic conjunctivitis caused by mountain

cedar pollen. Subjects were randomized to receive either loteprednol etabonate 0.2% ophthalmic suspension (LE) or placebo (vehicle), bilaterally, for 42 days.

Ocular safety evaluations included an external examination, slit lamp examination, tonometry and visual acuity taken prior to enrollment and at scheduled times during the study.

			Г	 		 		T
VISIT	Screen	1	2	3	4	•	•	Exit
(DAY)	-21 to	0	2/3	5-10	11-17	21-34	35-48	
PROCEDURE								
Informed Consent	Χp							
Inclusion/Exclusion_	ΧÞ							
Demographics, History	χb							
Medication History	ΧÞ							
Skin Test Results	Χp							
Pregnancy Test		Xª						χª
Visual Acuity		ΧÞ	×	x	×	x	x	
Ocular Signs & Symptoms		Χpε	×	х	х	x	x	
Intraocular Pressure		χb	×	x	x	х	x	
Undilated Fundus Exam		χb						×
Issue Medication		χÞ			х	×		
Issue Diary		Χþ						
Recover Diary				x	×	х		×
Investigator Global Assessment				x	х	х	x	
Recover Medications					x	x		×
Complete Exit Form	!			•				х
Dismiss Patient								×
Daily Environmental Allergen Counts ^d		×	x	x	x	x	x	×

^{*} Women of childbearing potential only.

^b Day -21 to Day-1 and Day 0 can be combined

Pre-treatment and 1 and 2 hours (± 10 min) post instillation of first drop

⁴ Daily Environmental Allergen Counts were required to be recorded until at least until 10 February 1996.

Inclusion Criteria:

- Adults, at least 18 years of age, of either sex and any race.
- Experience itching (at least 4+), and redness (at least 2+) due to pollen at Visit 1.
- Documentation of a positive allergy test to mountain cedar pollen by skin test within 12 months or RAST test within 36 months.

Exclusion Criteria

- Pregnant or lactating females.
- Females of childbearing potential who were not using adequate birth control.
- Previous allergic hypersensitivity to corticosteroid, loteprednol etabonate or to any component of the study medication.
- Expected concurrent ocular therapy with a non-steroidal anti-inflammatory agent, mast cell stabilizer, antihistamine, decongestant or beta-blocker during the period of masked medication treatment.
- Use of the medications listed above within 48 hours prior to Visit 1 (Day 0).
- Therapy with systemic or topical (ocular) corticosteroids within two weeks prior to the start of the study.
- Any abnormality preventing reliable applanation tonometry in either eye.
- Intraocular pressure that is greater than 21 mm Hg in either eye or any type of glaucoma.
- History of intraocular or laser surgery within the past six months.
- Best corrected (by pinhole) distance visual acuity (Snellen) in either eye worse than or equal to 20/100.
- Anticipated travel for more than a 24 hour period greater than 50 miles outside of the San Antonio/Austin area.
- Presence of any ocular pathology other than acute, seasonal allergic conjunctivitis (i.e., excluded is vernal conjunctivitis, GPC, viral or bacterial conjunctivitis or perennial allergic conjunctivitis).
- History of any severe/serious ocular pathology or medical condition (including systemic allergic disorders such as asthma or rhinitis) that could result in the patient's inability to complete this study.
- Previous participation in this study.
- Participation in any study under an IND within the past 30 days.
- Unlikely to comply with the protocol instructions for any reason (e.g., confusion, infirmity, alcohol or drug abuse).
- Contact lens wear during the course of the study.

Masking

While the physical appearance of the study medications was different (i.e., loteprednol etabonate 0.2% ophthalmic suspensions - opaque, white suspension; placebo (vehicle) - clear solution), this study was considered a double masked evaluation. Medications were supplied in opaque plastic containers with opaque dropper tips. Subjects were admonished not to discuss their medication with others on the study or in specific detail with the Investigator. The Investigator did not dispense study medication to subjects. A third party at the Investigator's office who was not responsible for patient assessments was given the responsibility of dispensing study medication to the subject, instilling medication when necessary and instructing the subject in study medication use.

Efficacy Criteria

The primary efficacy variables were bulbar conjunctival injection (sign) and ocular itching (symptom). Other supportive efficacy variables were discomfort, foreign body sensation, burning/stinging, photophobia, tearing and discharge (symptoms) and palpebral conjunctival injection, chemosis and erythema (signs).

An Investigator Global Assessment of the Control of Signs and Symptoms of SAC was recorded at Visits 3, 4, 5 and 6 for Days 0 to 7, 0 to 14 (inclusive), 15 to 28 and 29 to 42, respectively. This was to be based on the two previous clinical evaluations and daily diary data over a 14 day (approximate) period, except for Visit 3 (7 days). The rating after 2 weeks (Visit 4) was considered a secondary efficacy parameter.

Most signs-and symptoms were rated using a four point scale (0 - 3) where 0 = absent, 1 = mild, 2 = moderate and 3 = severe.

The Investigator Global Assessment used a 5 point scale (0-4) where 0=fully controlled, 1=reasonably controlled, 2=fairly controlled, 3=poorly controlled, 4=not controlled.

For selected parameters definitions were provided with the scales, as shown as follows:

Bul	bar Coniunctiv	<u>val Injection</u>
0	Absent	A normal, quiet eye; some subjects will exhibit rare vessels which are
1	Mild	naturally prominent either by location or a large normal vessel diameter. Slightly dilated blood vessels; color of vessels is typically pink; can be quadrantic (i.e., quadrant specific).
2	Moderate	More apparent dilation of blood vessels; vessel color is more intense (redder); involves the vast majority of the vessel bed.
3	Severe	Numerous and obvious dilated blood vessels; in the absence of chemosis the color is deep red - in the presence of chemosis, the leaking interstitial fluid may make the color appear less red or even pinkish; is not quadrantic.
ltchi	ing:	A sensation of the need to scratch or rub the eyelids or the periorbital area.
0	Absent	No desire to scratch or rub area.
1	Trace	Rare need to scratch or rub area but sensation is not completely absent.
2	Mild	Occasional need to scratch or rub area.
3	Moderate	Frequent need to scratch or rub area.
4	Severe	Constant need to scratch or rub area.

Disc	<u>charge</u> :	Involves the lash margin and adjacent eyelids and include crusts, collarettes, scaling, etc.
0	Absent	No abnormal discharge.
1	Mild	Small amount of mucopurulent or purulent discharge noted in the lower cul-de-sac. No true matting of eyelids upon awakening in the moming.
2	Moderate	Moderate amount of mucopurulent or purulent discharge is noted in the lower cul-de-sac. Frank matting together of eyelids in the morning upon awakening.
3	Severe	Profuse amount of mucopurulent or purulent discharge noted in the lower cul-de-sac and in the marginal tear strip.
Pho	tophobia:	Abnormal ocular or periocular discomfort, pain or sensitivity upon exposure to light.
0	Absent	Absence of positive sensation
1	Mild	Very minimal light intolerance which may require some degree of sunglass protection to eliminate the symptom, notice primarily in sunlight.
2	Moderate	Infrequent or intermittent discomfort in the globe associated with exposure to room light or sunlight which is only partially relieved by dark glasses or subdued light. The symptoms still persist to some degree even with sunglasses.
3	Severe	Constant or nearly constant exquisite pain in the eye that is not relieved by sunglasses and is only relieved by total occlusion of the eye. This total occlusion can be achieved with an eye patch or by closing the eyes. This sensation is so significant that frequently bed rest and occasionally systemic sedation is required to relieve this severe grade of symptom.
hy Crit	eria	

Safety Criteria

Ocular safety examinations included an external examination, slit lamp examination, funduscopy, applanation tonometry and visual acuity, taken prior to enrollment and at scheduled times during the study. Systemic safety evaluation was obtained by subject comment with physician follow-up. Safety parameters were tabulated to identify those showing a difference in incidence rate between treatment groups.

Allergen Counts

During the study all Investigators were required to record local environmental allergen counts from the time of the first screening visit until all patients had completed the study.

The study was carried out during the mountain cedar pollen season in South Central Texas (December 1995 to February 1996). For safety evaluation some patients continued taking the test article beyond the active pollen season. The final on-study patient day was 9 March 1996.

Concurrent Therapy

The following systemic medications were allowed to be used concurrently: NSAID's, oral birth control pills, estrogen replacement, thyroid preparations, insulin, hypoglycemic agents and anti-microbials for non-ocular conditions. The following nasal and ocular rescue medications could be used as needed to control EXCESSIVE nasal and ocular allergic symptoms: phenylephrine hydrochloride nasal solution (Neo-Synephrine®, Dristan®, etc.) and cromolyn sodium nasal solution (Nasalcrom®) could be used at the onset of nasal allergic symptoms. The only allowed ocular rescue medication was artificial tears. Those individuals receiving immunotherapy (allergy shots) should have been on a stable regimen prior to the last allergy season and must have no unusual changes to their dosing regimen during the period of masked medication and within two (2) days prior to the Enrollment exam. Patients receiving concurrent medication during the study that was prohibited were to be discontinued from the study as a protocol violation. Patients requiring additional ocular medication other than the masked study medication or artificial tears during the study were to be discontinued from the study as a protocol violation or a treatment failure and placed on appropriate medication. Specifically excluded were: ocular steroids, ocular non-steroidal anti-inflammatory agents, and ocular mast cell stabilizers; systemic steroids, systemic antihistamines and systemic decongestants. All concurrent drug use was to be documented in the Case Report Form.

The target sample size was 64 evaluable patients per treatment group (total = 128 patients). There were 133 patients randomized to treatment out of 387 patients screened. Sixty six were assigned to receive LE and 67 were assigned to receive placebo. The first patient was enrolled on 19 December 1995 and the last patient visit occurred on 9 March 1996. Patients who discontinued treatment before Visit 6 (Day 42) were considered to have not completed the study. One hundred twenty six (126) patients completed treatment through Visit 6, and thus, the study. Four (4) patients (2 on LE; 2 on placebo) were discontinued due to a medical event, one patient was lost to follow-up (placebo), and two patients (both on placebo) due to reasons unrelated to the study as shown below:

•	474 0004 (5)		The set of the second second
•	174-3004 (Placebo)	Day 16	Terminated: Severe itching
•	174-3014 (Placebo)	Day 7	Discontinued: Protocol violation
			(Patient used Alka-Seltzer Plus®)
•	174-3019 (Placebo)	Day 22	Discontinued: Needed to travel
•	175-3084 (LE)	Day 7	Terminated: Elevated IOP, O.U.
•	175-3093 (LE)	Day 31	Terminated: Acute pharyngeal reaction, headache
•	175-3097 (Piacebo)	Day 1	Terminated: Viral conjunctivitis
•	178-3170 (Placebo)	Day 0	Discontinued: Lost to follow-up

There was one patient with no on-treatment evaluation (178:3170, placebo). This patient was <u>not</u> included in the intent to treat analysis. For the intent to treat analysis, all patient data for Visit 2 (Day 2/3), Visit 3 (Day 7) and Visit 4 (Day 28) were included in the primary intent to treat analysis without regard to whether the visits were in the day range specified in the protocol.

A threshold pollen count of 100/m³ was set a priori in the study protocol. Pollen counts by date for each of the two cities with investigational sites are shown. For the efficacy analysis, no visits within the first two weeks were disqualified due to evaluations after the allergy season. For visits 5 and 6 (four and six weeks), only visits which were in the defined allergy season were used in the intent-to-treat efficacy analysis.

The pollen count in Austin (Investigator 174) was over 100/m³ after 11 December 1995 and the first patient entered the study on 21 December 1995. The pollen count in San Antonio (Investigators 175 and 178) was over 100/m³ after 14 December 1995 and the first patient entered on 19 December 1995. All patients had itching and bulbar injection of sufficient severity to qualify.

After 28 January 1996 in Austin and 9 February 1996 in San Antonio pollen counts were mostly under 100/m³ and these dates were determined to be the end of the mountain cedar season prior to unmasking the study. There were 3 patients with Visits 3 and 4 after the season and 12 patients with Visit 4 afterwards.

	Austin	San Antonio
Investigator(s)	174	175, 178
Pollen > 100/m ³	12 December 1995	15 December 1995
First patient enrolled	21 December 1995	19 December 1995
Last patient enrolled	25 January 1996	27 January 1996
Poilen < 100/m³	28 January 1996	9 February 1996

A valid visit for a patient required that the patient had to take study medication within 48 hours. There were 2 scheduled visits that occurred more than 48 hours after the last dose; these were Visit 6 evaluations that occurred after the end of the mountain cedar season. A tighter criterion was applied for a per protocol valid visit. Study medication had to be taken within 4 hours of the visit, disallowed medications were not to be taken prior to the visit and the visit had to be within the day range specified in the protocol.

One LE and two placebo patients exceeded the four hour limit at their final visit, but these visits were after the end of the mountain cedar season. Disallowed medications were taken by 2 LE and 2 placebo patients. One placebo patient (174:3014) was dropped because of this deviation from the protocol; one (175:3093) was discontinued at an unscheduled visit after being instructed to begin treatment with a disallowed medication. The LE patient (174:3016) and the placebo patient (178:3189) continued in the study after their deviations. There were 3 LE patients and 1 placebo patient who were off-schedule for visits that occurred after the end of the mountain cedar season. There were 1 LE (178:3182) and 2 placebo patients (175:3116 and 178:3184) who were off schedule during the season for Visit 3, 4 and 2, respectively.

It was anticipated that the mountain cedar season would end before Visits 5 (Day 28) and Visit 6 (Day 42) for many patients. These visits were not to be included in the intent to treat statistical analysis of Visit 5 or 6. The per protocol criteria disallowed any visit after the end of the mountain cedar season.

There was 1 placebo patient (175:3097) who developed viral conjunctivitis on Day 1 in the right eye. The ratings for the left eye were used in the intent to treat analysis, but failed to meet the criteria of an on-schedule visit for per protocol. There were 2 additional missed visits (LE 174:3009 Visit 3 and placebo 174:3025 Visit 2).

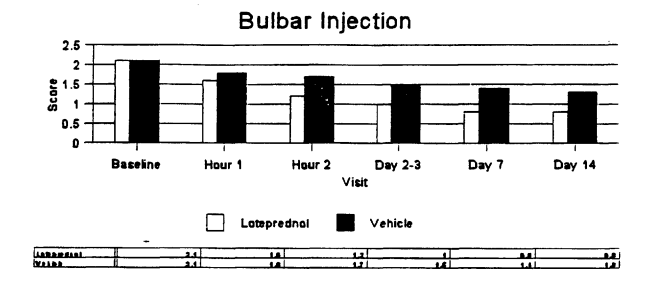
APPEARS THIS WAY

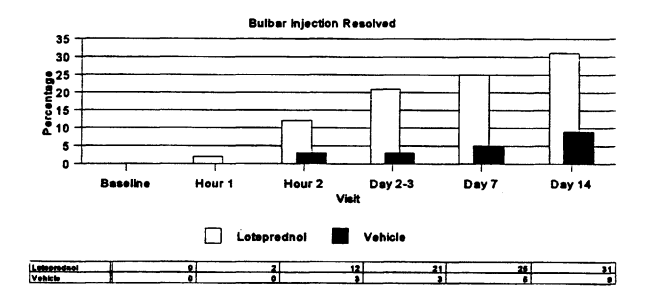
Pollen Counts

WEEK								
BEGINNING	CITY	MONDAY	TUESDAY	WEDNESDAY	THURSDAY	FRIDAY	SATURDAY	SUNDAY
11DEC95	AUSTIN	47	1720	4005	3750	3250	325	300
	SAN ANTONIO	0	0	20	0	990	14100	1590
18DEC95	AUSTIN	188	287	105	253	1700	355	25
	SAN ANTONIO	1870	2330	830	100	70	50	90
250EC95	AUSTIN	70	1800	4515	8000	1145	2020	3010
	SAN ANTONIO	50	160	40	16500	7250		
01JAN96	AUSTIN	3002	2960	1150	3885	3650	3240	1950
	SAN ANTONIO	50000	18100	1420	700	200	12200	13480
08JAN96	AUSTIN	1980	1460	2085	1500	2350	2100	2025
	SAN ANTONIO	_ 25200	650	5400	29000	21500	50000	44000
15JAN96	AUSTIN	2285	1950	1900	2400	1435	1645	3545
	OINOTHA HAZ	11400	50000	6500	19000	5220	810	3200
22JAN96	AUSTIN	875	1520	1655	425	330	600	180
	SAN ANTONIO	360	8000	4220	200	150	500	370
29JAN96	AUSTIN	45	40	25	80	20	20	40
	SAN ANTONIO	260	100	9 70	160	0	50	0
05FEB96	AUSTIN	135	25	260	45	20	25	18
	SAN ANTONIO	0	1220	1570	250	200	0	0
12FEB96	AUSTIN	18	0	0	0	0	0	0
	SAN ANTONIO	0	0	0	60	320	0	0

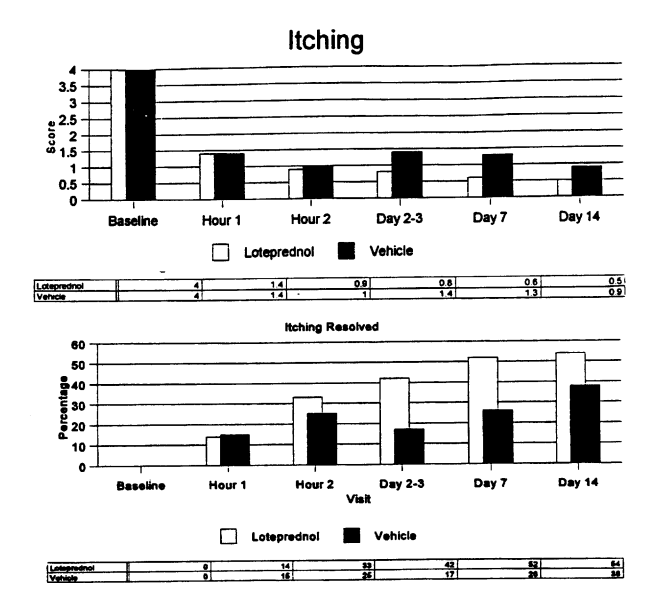
APPEARS THIS WAY ON ORIGINAL

_					Reason: Study Incomplete					
Treatment		and	omized Con S	npieted tudy	Lack of Efficacy	Adve Eve		st to Other w-up Unrelated		
LE	66	,	64 (9	7%)	0 (0%)	2 (3%)	0 (0%)	0 (0%)		
Placebo	67	•	62 (9	3%)	0 (0%)	2 (3%)	1 (1%)	2 (3%)		
			LOTEPREDNOL		INV 174					
age	N		66	67	34	60	39			
		AN	40.6	41.5	41.6	42.1	39.0			
		IN Ax	-							
GENDER								•		
MALE	N	*	33 50%	32 48%	14 419					
FEMALE	N	*	33 50%	35 52%	20 599		0% 18 46%			
			HO:LE=PL	p=0.863	HO: INV	EQUAL p=	0.543			
RACE										
CAUCASIAN	N	*	42 64%	41 61%	30 889					
HISPANIC	N	*	22 33%	19 28%	2 6		0% 21 54%			
OTHER	N	*	2 3%	7 10%	2 69		3% 5 13%			
			HO:LE=PL	p=0.223	HO:INV	EQUAL p=	0.001			
IRIS	•									
LIGHT	N	*	29 44%	26 39%	19 568		0% 12 31%			
DARK	N	X	37 56%	41 61%	15 449	_				
			HO:LE=PL	p=0.599	HO: INV	EQUAL p=	0.090			
CITY										
AUSTIN	N	*	16 24%	18 27%	34 100		0% 0 0%			
SAN ANTONIO	N	¥	50 76%	49 73%	0 0					
			HO:LE=PL	p=0.843	HU:1NV	EQUAL p=	0.001			
DACEL THE DOLL	CN.			•						
BASELINE POLL			64	66	34	58	38			
	N	AN		1164.5	2154.1	12185.5	17126.6			
	SC			5056.8	1010.3	16284.0	19966.3			
	M]		100/0.0 1		1010.0	_V_V_	2200.0			
	M									
0-500	N	*	16 25%	15 23%	4 12%	16 28	11 29%			
501-1000	N	ž	9 14%	9 14%	0 0%	14 24				
1001-5000	N	ž	18 28%	19 29%	30 88%	4 7				
5001-15000	N	ž	5 8%	5 8%	0 0%	5 9		•		
>15000	N	ž	16 25%	18 27%	0 0%	19 33				
- 10000	••	-4	HO:LE=PL :	¥	-	QUAL p=0				





A higher percentage of patients in the loteprednol group had resolution of redness compared to the vehicle group. The means scores are not impressively different.



A higher percentage of patients in the loteprednol group had resolution of itching compared to the vehicle group. The means scores are not impressively different.

Bulbar Inje	ection	OBSERVED	RATINGS:	MEAN OF EVES (C	n-051/2			
		IT 1 (DAY D)		VISIT 2	VISIT 3	VISIT 4	VISIT 5	VISIT 6
	BASELINE	HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42
LOTEPREDNOL (LE)					•	5 ,4	UP11 (20)	2011 42
DISTRIBUTION	N 3	H 1	N 1	N 1	H 1	N 3	N 1	N 2
0:ABSENT	0 01	1 23	8 123	14 212	16 251	20 31%	6 223	1 132
0.5-1:HILD	0 03	20 30%	36 552	35 532	40 623	26 55%	17 633	5 631
1.5-2:HODERATE	56 853	40 612	20 301	17 268	9 14%	9 148	4 158	2 25\$
2.5-3:SEVERE	10 15%	5 81	2 33	0 0%	0 03	0 0%	0 03	0 Ot
Ħ	66	66	66	66	65	65	27	8
HEAN	2.1	1.6	1.2	1.0	0.8	0.8	0.9	1.1
STANDARD ERROR	0.0	n i	0 1	0.1	0.3	ถ 1	6.1	0.8
MIN. MAX								
MEDIAN	2.0	2.0	1.0	1.0	1.0	1.0	1.0	1.0
0- 40700 - 40- 1								
PLACEBO (PL)								
DISTRIBUTION	H S	# 1	N 3	H \$	N 3	N 3	N 2	N 3
O:ABSENT	0 03	0 03	2 33	2 31	3 5%	6 91	2 82	0 02
0.5-1:MILD 1.5-2:MODERATE	0 02	13 19%	15 22%	26 401	28 433	33 524	7 27%	4 50%
	54 81%	40 723	46 723	34 523	33 513	23 361	16 623	3 381
2.5-3:SEVERE	13 194 67	6 91	2 31	3 5%	1 23	2 31	1 43	1 134
MEAR	2.1	67 1.8	67	66	65	64	26	8
STANDARD ERROR	0.0	1.B	1.7	1.5	1.4	1.3	1.5	1.6
HIN HAX	U 11 ·		11 1	A 1	A 1	Ωı	n ı	u s
MEDIAN	2.0	-						
· CDI-M	2.0	2.0	2.0	1.5	1.5	1.8	1.5	1.5
				CHANGE FROM BAS	C) THE /006	COUCO BACE	. 195 6-31	
LOTEPREDNOL (LE)	•	HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42
FREQUENCY DISTRIBUTE	ON	H E	N 1	N 1	N I	N E	N 1	# 1
IMPROVED -3		î 23	1 23	î 23	1 23	îzî	0 03	0 03
-2.52		0 003	8 123	13 203	17 283	20 313	6 223	1 133
-1.51		22 333	35 533	35 533	39 601	36 55%	17 63%	5 633
UNCHANGED -0.5 - 0.	5	43 65k	21 323	17 263	8 123	8 123	4 15%	2 25%
1.5 - 1		0 02	1 23	0 0%	0 03	0 03	0 0%	0.03
MORSENED 2.5 - 2		0 Ot	0 01	o ox	0 03	0 01	0 03	0 02
								J (4
N	66	66	66	66	65	66	27	A
MEAN	2.1	-0.5	-0.9	-1.1	-1.3	-1.3	-1.2	-0.9
STANDARD ERROR	0.0	n t	A 1	0 1	0.1	0.1	0.1	0.2
MJN, MAX								
HEDIAN	∠. U	-U. 5	-1.0	- L. Li	-1.U	-1.0	-1.0	-1.0
								•••
PLACEBO (PL)								
FREQUENCY DISTRIBUTION	ON	H I	x 1	N I	N I	N S	N \$	N I
IMPROVED -3		0 03	0 0%	0 02	2 33	1 23	0 02	0 01
·2.5 · ·2		1 13	3 43	5 81	3 52	8 131	4 15%	0 01
-1.51	_	17 25%	20 30 1	29 45%	32 493	33 523	8 313	4 50%
UNICHANGED -0.5 - 0.5	5	49 73%	44 662	30 462	Z7 4ZX	21 333	14 542	4 50%
1.5 - 1		0 03	0 04	1 23	1 23	1 23	0 0%	0 02
NORSENED 2.5 - 2	_	0 03	0 0%	0 03	0 02	0 03	0 01	0 03
•	ଟ	67	ा	65	65	64	26	8
MEAN STANDARD ERROR	2.1	-0.3 8 1	-0.4	-0.7	•0.7	-0.9	-0.7	-0.5
MIN. MAX	u 11	B 1	8 1	n۱	8 3	0 1	0 1	0.7
MEDIAN	2.U	0.0						
HEUTH	2.0	V. U	U.U	-1.U	-1.V	-1.4	-U.5	-U.5
UNIVARIATE ANALYSES	BASEL INE	HOUR 1	HOUR 2	VISIT 2	VISIT 3	VISIT 4	VISIT 5	WICHT &
INVEST p-VALUE(c)	0.792			VIJI. E	-1911 3	41311 4	41211.2	VISIT 6
TRT p-VALUE(d)	0.265	0.274	0.000	0.001	0.000	0.006	0.030	0.135
TREATMENT EFFECT[e]	0.0	0.0	-0.5	-0.5	-0.5	-0.5	-0.5	0.0
95% CONF LINITS	0.0. 0.0	0.0. 0.0 -	1.00.5	-0.5. 0.0 -	1.00 5	-04.00	-1000	-1000
(a) OBSERVED RATINGS A	WE THE HEAR	OF BOTH EYES	AT THE Y	ISIT: CHANGE FROM	BASELINE	IS OBSERVED	RATING - MASS	LINE RATING
THUS, IMPROVENENT	IS A NEGATIVE	E NUMBER						

Visits 5 and 6 do not have sufficient numbers of patients for evaluations of efficacy.

⁽a) DISEATED BATTIMES ARE THE MEAR OF BUTH EYES AT THE \$1331;CHARME FROM BASELINE TO MISSEATED INCLINE TO MISSEATED INCLINE TRUTH THE THEOLOGY.

THIS, IMPROVEMENT IS A DEGATIVE NUMBER

(b) REPEATED DEASURES AMALYSIS OF CONARIANCE WITH AN ESTIMATE OF OVERALL TREATMENT EFFECT (LE-PL): NEGATIVE TREATMENT EFFECTS INDICATE LOTERPREDUCL FAVORED OVER PLACEBO

(c) COCHRAM-MARTEL-MEISZEL TEST FOR EQUALITY OF ROM (THEATMENT) MEAN MARKS AT BASELINE

(d) COCHRAM-MARTEL-MEISZEL TEST FOR EQUALITY OF ROM (THEATMENT) MEAN MARKS CONTROLLING FOR INVESTIGATOR

(e) LARGE SAMPLE ESTIMATE OF THE MEDIAN DIFFERENCE BETWEEN TREATMENT GROUPS (LE-PL) AND ITS CONFIDENCE LIMITS

Itching			OBSERVED RAT	INGS: MEAN (OF EYES (OD-	-051/2		
•	VISI	(DAY 0)		VISIT 2	VISIT 3	VISIT 4	VISIT 5	VISIT 6
	BASEL INE	HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42
LOTEPREDUCL (LE)	· N 2	N 3	H I	N E	H 2	N \$	N 3	N \$
O:ABSENT	0 03	9 148	22 333	28 423	34 52%	35 541	20 748	6 752
0.5-1:TRACE	0 03	25, 423	29 443	22 423	23 351	25 381	3 118	2 251
1.5-2:MILD	0 0%	20 301	12 18%	7 118	7 118	3 58	3 118	0 0%
2.5-3:HODERATE	0 02	7 11%	3 5%	2 31	1 2% 0 0%	2 3% 0 0%	1 4% C 0%	0 0%
3.5-4:SEVERE	661003	2 33 66	p 01. 66	1 2% 66	65 65	66	27	8
N NEAN	66 4.0	1.4	0.9	0.8	0.6	0.5	0.4	0.3
STANDARD ERROR	0.0	0.1	0.1	0.1	0.1	n 1	0.3	^ •
MIN. MAX								
HEDIAN	4.U	1.0	1.0	1.0	0.0	0 0	0.0	0.0
PLACEBO (PL)								
DISTRIBUTION	N 1	N \$	N S	N X	N \$	N 3	N 3	N 1
O:ABSENT	0 0%	10 153	17 258	11 17%	17 26%	24 384	18 693	21003
0.5-1:TRACE	0 93	30 45%	29 431	22 341	24 37%	25 391	2 81	0 03
1.5-2:MILD	0 0%	16 24%	19 28%	22 348	13 20%	10 163	4 15% 2 8%	0 0%
2.5-3:MOCERATE	0 0%	11 164	2 33 0 04	9 14k 1 2k	8 127 3 58	4 6% 1 2%	2 0a 0 03	0 03
3.5-4:SEVERE	67100% 67	0 0% 67	0 UA 67	66	65 .	· 64	26	8
MEAK	4.0	1.4	1.0	1.4	1.3	0.9	0.6	0.0
STANDARD ERROR	2.0	A 1	A 1	A 1	۸ ۱	۸ ۱	4 3	n n .
MIN, MAX								
MEDIAN	4.0	1.0	1.0	1.0	1.0	1.0	g.C	0.0
			CHA	NGE FROM BAS	SELINE (OBS	ERVED - BASE		
		VISIT 1		VISIT Z	VISIT 3	VISIT 4	VISIT 5	VISIT 6
LOTEPREDMOL (LE)		HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42 N I
FREQUENCY DISTRIBUTION	l	N \$	N S	N I 20 423	N 2 34 523	N % 35 54%	20 74%	6 753
IMPROVED -4		9 14\$ 28 42\$	22 338 29 441	28 423	23 35%	25 381	3 113	2 258
-3.5 - ·3 -2.5 · -2		20 30%	12 183	7 113	7 11%	3 5%	3 11%	0 08
-1.51		7 118	3 5%	2 33	1 23	2 33	1 43	0 0%
UNCHANGED -0.5 - 0.5		2 32	0 0%	1 23	0 0%	0 02	0 0%	0 0%
N	66	5 6	66	66	66	65	27	8
HEAK	4.0	-2.6	-3.1 n 1	-3.2 0.1	-3.4 n 1	-3.5 n 1	-3.6 n 2	-3.8 0.2
STANDARD ERROR	4.0	n 1	n v		" '		" /	,
MIN, MAX MEDIAN	4.0	-3.0	-3.0	-3.0	-4.0	-4.0	-4.U	-4.U
MEDINA	4.0	-0.0	-5.0	•.•				
PLACEBO (PL)								N 2
FREQUENCY DISTRIBUTION	i	N T 10 15%	N \$ 17 25%	N X 11 17X	N % 17 26%	N 3 24 384	N 1 18 691	81003
IMPROVED -4 -3.5 · -3		30 458	29 43\$	22 343	24 37%	25 391	2 84	0 03
·2.5 · ·2		16 243	19 253	22 343	13 203	10 162	4 15%	0 03
-1.51		11 163	2 38	9 143	8 123	4 6%	2 81	0 01
UNCHANGED -0.5 - 0.5		0 0%	0 0%	1 23	3 58	1 24	0 0%	0 02
K	67	67	67	65	65	64	26 ∙3,4	8 -4.0
MEAN	4.0	-2.6 n 1	-3.0 n 1	-2.6 n 1	-2.7 0.1	-3.1 n 1	*3. * n ?	-4.0
STANDARD ERROR	a a	n 1			•••	** .		
MIN. MAX MEDIAN	4.0	-J.U	-3.0	-J.U	-3.0	-3.0	-4.0	-4.0
				_	.62			
TREATMENT EFFECT		-0.02 -0.29, 0.3	M		0.37			
958 CONF LIMITS SUPPORTIVE		-J.EF. V.	•	- 4 , 00				
UNIVARIATE ANALYSES		HOUR 1	110UR 2	VISIT 2	VISIT 3	VISIT 4	VISIT S	9 TIZIV
TRT p-VALUE[c]		0.590	0.304	0.000	0.000	0.034	1.000	0.134
TREATMENT EFFECT(d)		0.0	0.0	-1.0	-1.0	0.0	0.0	0.0
95% COMF LIMITS		0.0, 0.0	0.0, 0.0 -1	5	-1.0. 0.0	-0.5, 0.0	0.0, 0.0	0.0, 1.0

Visits 5 and 6 do not have sufficient numbers of patients for evaluations of efficacy.

[[]a] OBSERVED RATINGS ARE THE HEAM OF BOTH EYES AT THE VISIT. BASELINE ITCHING WAS SEVERE (4) FOR ALL PATIENTS.

CHANGE FROM BASELINE IS OBSERVED RATING - 4: THUS, IMPROVEMENT IS A NEGATIVE NUMBER

(b) REPEATED MEASURES AMALTSIS OF VARIANCE WITH AN ESTIMATE OF OVERALL TREATMENT EYFECT (LE-PL): NEGATIVE TREATMENT EFFECTS INDICATE LOTEPREDUCL FAVORED OVER PLACEBO.

(c) COCHRAM-HAMTEL-HAENSZEL TEST FOR EQUALITY OF ROM (TREATMENT) MEAN RAMES CONTROLLING FOR INVESTIGATOR

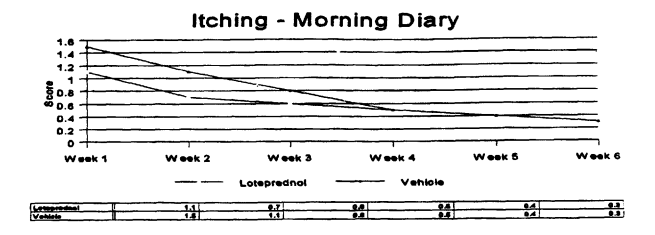
(d) LARGE SAMPLE ESTIMATE OF THE MEDIAN DIFFERENCE BETMEEN TREATMENT GROUPS (LE-PL) AND ITS COMFIDENCE LIMITS

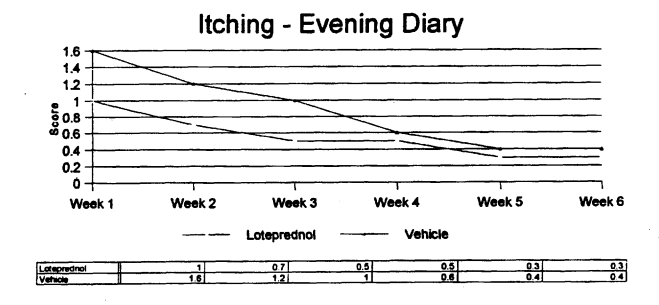
							State 4
Measure:	Treatment	N	Hour 1	Hour 2	Day 2/3	Day 7	Day 14
Discomfort:	LE	65	14%	35%	43%	47%	66%
	Placebo	66	17%	27%	22%	41%	52%
Foreign body sensation:	LE	50	40%	48%	57%	68%	84%
	Placebo	62	35%	52%	43%	60%	61%
Burning/Stinging:	LE	64	33%	36%	53%	60%	68%
	Placebo	67	30%	33%	38%	43%	53%
Photophobia:	LE	56	48%	52%	64%	64%	65% ⁻
	Placebo	53	43%	49%	52%	55%	54%
Tearing: -	LE	64	41%	·50%	61%	67%	70%
	Placebo	64	34%	44%	42%	58%	57%
Discharge:	LE	52	58%	71%	63%	63%	63%
	Placebo	44	80%	80%	56%	67%	71%
Palpebral injection:	LE	66	2%	9%	23%	25%	29%
	Placebo	67	0%	3%	6%	8%	13%
Chemosis:	LE	63	10%	16%	33%	35%	35%
	Placebo	60	2%	5%	24%	22%	22%
Erythema:	LE	61	13%	25%	49%	51%	45%
	Placebo	59	10%	17%	28%	28%	36%

N is the intent-to-treat sample size of patients with the sign or symptom present at baseline.

Reviewer's Comments:

With the exception of the measure of Discharge, the loteprednol group almost always had a higher percentage of patients with resolved signs and symptoms.





The graphs show overall improvement from baseline in both groups and little difference between groups.

Intraocular Pressure Elevations

dan ini ana			androment States	and second	
LE	> 15	0	0	0	0
	10-15	0	0	0	0
	6-9	6	6	8	4
	< 6	59	59	57	60
Placebo	> 15	0	0	0	0
	10-15	0	0	0	0
	6-9	0	4	1	1
	< 6	65	60	61	61

The distributions (above and below) displayed are the changes in IOP in the eye with the greatest increase from baseline in IOP

			OBSERVED		
		VISIT 3	VISIT 4	VISIT 5	VISIT 6
LOTEPREDNOL	BASELINE	DAY 7	DAY 14	DAY 28	DAY 42
DISTRIBUTION	N I	N \$	N \$	N X	N \$
< 20 MM HG	66100\$	58 89%	59 91%	58 89\$	57 89%
20 - 25 MM HG	0 0%	6 9	6 9%	7 11%	7 11%
26 - 31 MM HG	0 0%	1 2%	0 0%	0 0%	0 0%
> 31 MM HG	0 02	0 0%	0 0%	0 0%	0 0%
N	66	65	6 5	65	64
MEAN	14.6	16.1	15.8	16.1	15.7
STANDARD ERROR MIN. MAX	0.3	0.4	0.4	0.4	0.4
MEDIAN	14.5	16.0	16.0	16.0	15.0
PLACEBO (PL)					
< 20 MM HG	65 97*	64 98%	63 98%	59 95%	58 94%
20 - 25 MM HG	2 3%	1 23	1 2%	3 5%	4 63
26 - 31 MM HG	0 0%	0 03	0 0%	0 0%	0 0%
> 31 MM HG	0 0%	0 02	0 02	0 0%	0 0%
N	67	65	64	62	62
MEAN	14.4	14.5	14.7	15.0	15.1
STANDARD ERROR	0.3	0.4	0.4	0.4	0.4
MIN. MAX MEDIAN	14.0	14.0	15.0	14.5	15.3

Reviewer's Comments:

Elevations in IOP were seen more frequently in the loteprednol group.

Adverse Experiences: (>2%)

SPECIAL SENSES	PATIENTS - PATIENTS REPORTING EVENT			TOTAL	SEVERITY OF EVENTS		
	= PATTENTS AT RISK		ING EVENT	MUMBER Of Events	MILD	MODERATE	CENEDE
BODY AS A WHOLE -Any Event	NI VION	A1 LD	DI ONCE	O. EVENIS	mich	HODEIVIL	JETERE
LOTEPREDNOL	66	21	323	24	12	9	3
PLACEBO	67	20	301	23	16	ž	Õ
CHEMOSIS (EYE/CONJ)						•	•
LOTEPREDNOL	66	11	172	14	13	1	0
PLACEBO	67	15	221	17	16	1	0
ITCHING. EYE (EYE/GEN)							
LOTEPREDNOL	66	10	15%	10	9	1	0
PLACEBO	67	25	372	28	19	7	2
HEADACHE (HEAD)							
LOTEPREDNOL _	66	10	151	11	5	4	2
PLACEBO	67	10	152	10	9	1	0
ERYTHEMA. EYELIDS (EYE/API	P)						
LOTEPREDHOL	66	7	112	7	7	0	0
PLACEBO	67	6	9t	7.	4	3	0
FLU SYNDROME (GEN)							
LOTEPREDNOL	66	6	91	6	1	4	1
PLACEBO	67	0	02	0	0	D	0
BURNING/STINGING, EYE, NOT	ON INSTIL	LATION (EYE/GEN)				
LOTEPREDNOL	66	6	91	6	5	1	0
PLACEBO	67	6	91	9	3	3	3
DISCHARGE, EYE (EYE/GEN)							
LOTEPREDNOL	66	6	91	6	6	0	0
PLACEBO	6 7	17	251	20	17	1	2
EPIPHORA (EYE/APP)							
LOTEPREDNOL	66	5	82	5	4	0	1
PLACEBO	67	14	213	14	7	6	1
FOREIGN BODY SENSATION (EY	E/GEN)						
LOTEPREDNOL	66	5	81	5	4	1	0
PLACEBO	67	11	16\$	12	8	2	2
DRY EYES (EYE/GEN)							
LOTEPREDNOL	66	4	61	5	3	2	0
PLACEBO	67	2	31	2	1	1	0
DISCOMFORT. EYE (EYE/GEN)							
LOTEPREDNOL	66	4	61	4	4	0	0
PLACEBO	67	3	41	3	2	0	1
INJECTION (EYE/CON)							
LOTEPREDNOL	66	3	51	3	2	1	0
PLACEBO	67	16	243	18	12	4	2
BURN/STING. EYE. ON INSTILL							
LOTEPREDNOL	66	3	51	3	2	1	0
PLACEBO	67	4	61	4	2	1	0

NDA 20-803: loteprednol etabonate ophthalmic suspension, 0.2%

EYE PAIN (EYE/GEN)							
LOTEPREDNOL	66	3	51	3	3	0	0
PLACEBO	67	1	1\$	1	1	Q	0
ALLERGIC REACTION (GEN)							
LOTEPREDNOL	66	2	32	2	2	0	0
PAIN (GEN)			•				
LOTEPREDNOL	66	2	31	2	2	0	0
PLACEBO	67	1	12	1	0	1	0
INFECTION. EAR. NOS (EAR	R/GEN)						
LOTEPREDNOL	66	2	31	2	0	2	0
PLACEBO	67	0	02	0	0	0	0

Reviewer's Summary of Safety and Efficacy

Marginal efficacy has been demonstrated in the resolution of itching and redness. Adverse experiences in this limited study (42 days) were generally confined to mild to moderate ocular events. There was an increased chance of increased IOP during use.

8.1.2 Study #2 Protocol: 144 Title: Safety and efficacy of loteprednol etabonate in the treatment of seasonal allergic conjunctivitis (QID) dosing. Objective: To evaluate the efficacy and safety of loteprednol etabonate 0.2% ophthalmic suspension in the treatment of signs and symptoms of environmental seasonal allergic conjunctivitis. Study Design: A randomized, double-masked, placebo controlled, parallel group multicenter (4 sites) study. Population: There were 135 adult patients, exhibiting signs and symptoms of environmental seasonal allergic conjunctivitis, coincident with elevated levels of an airborne pollen to which they had a demonstrated skin prick or RAST reaction.

Test Drug Schedule: Same as Study #2 (Protocol #143)

Number of Patients Enrolled Investigators: Richard B. Briggs, M.D. (#181) 30 Brackenridge Professional Building 1313 Red River, Suite 206 Austin, TX 78701 Larry L. Lothringer, M.D. (#179) 36 The Center for Corrective Eye Surgery 303 East Quincy San Antonio, TX 78215 Jay M. Rubin, M.D. (#180) 32 Eye Physicians 999 E. Basse Road San Antonio, TX 78220 37 David G. Shulman, M.D. (#176) Eye Clinic 999 E. Basse Road, Suite 116

San Antonio, TX 78220

Study Plan: Same as Study #2
Inclusion Criteria: Same as Study #2
Exclusion Criteria: Same as Study #2
Masking: Same as Study #2

Efficacy Criteria: Concurrent Therapy: Same as Study #2 Same as Study #2

Patient Disposition:

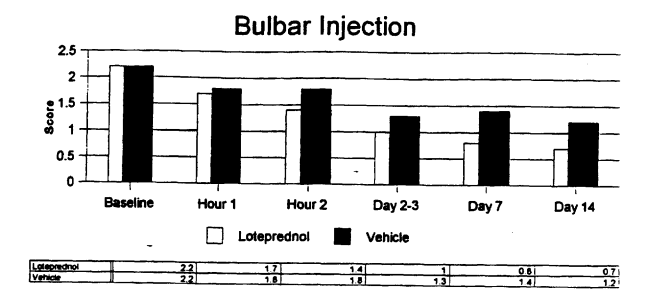
The target sample size was 64 evaluable patients per treatment group (total = 128 patients). There were 135 patients randomized to treatment out of 480 patients screened. Sixty seven were assigned to receive LE and 68 were assigned to receive placebo. The first patient was enrolled on 18 December 1995 and the last patient visit occurred on 9 March 1996. Patients who discontinued treatment before Visit 6 (Day 42) were considered to have not completed the study. One hundred twenty eight (128) patients completed treatment through Visit 6. Three (3) patients (1 on LE; 2 on placebo) were discontinued due to a medical event, 3 (1 on LE; 2 on placebo) were lost to follow-up and 1 patient (LE) due to reasons unrelated to the study as shown below:

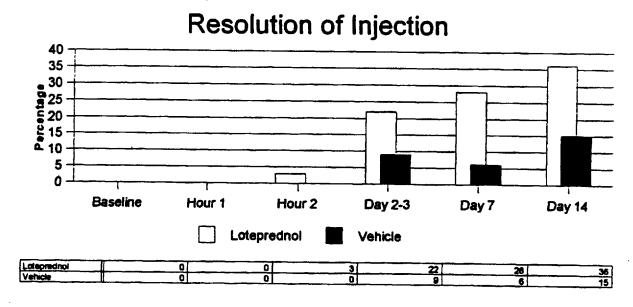
	Delow.	
•	176:4013 (placebo) Day 0	Lost to follow-up
•	176:4016 (placebo) Day 14	Increased IOP
•	179:4081 (LE) Day 2	Diagnosed with ovarian tumor - scheduled for
		surgery
•	179:4100 (LE) Day 0	Lost to follow-up
•	180:4135 (placebo) Day 6	Eye spasm (OD) upon instillation
•	180:4158 (LE) Day 7	Hospitalized following motor vehicle accident
•	180:4159 (placebo) Day 28	Lost to follow-up

	Austin	San Antonio
Investigator(s)	181	176, 179,180
Pollen > 100/m³	12 December 1995	15 December 1995
First patient enrolled	18 December 1995	18 December 1995
Last patient enrolled	26 January 1996	27 January 1996
Polien < 100/m³	28 January 1996	9 February 1996

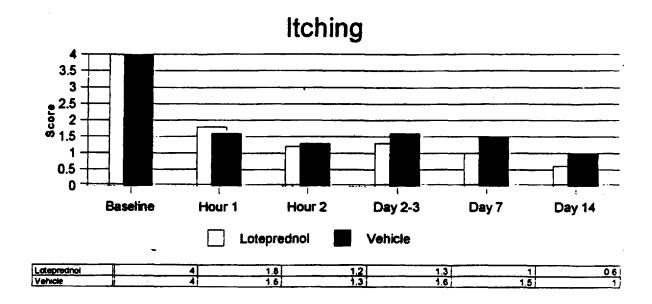
WEEK BEGINNING	CITY	MONDAY	DIECDAY	HERMECOAV	THURSDAY	FRIDAY	£470mnav	CIMPAY
DCO11M11M2	CITT	MUNUAY	IUESDAT	WEDNESDAY	INUKSUAT	PRIDAT	SATURDAY	SUNDAY
11DEC95	AUSTIN	47	1720	4005	3750	3250	325	300
	SAN ANTONIO	0	0	20	0	990	14100	1590
18DEC95	AUSTIN	188	287	105	253	1700	355	25
	SAN ANTONIO	1870	2330	830	100	70	50	90
25DEC95	AUSTIN	70	1800	4515	8000	1145	2020	3010
	SAN ANTONIO	50	160	40	16500	7250		
01JAN96	AUSTIN	3002	2960	1150	3885	3650	3240	1950
	SAN ANTONIO	50000	18100	1420	700	200	12200	13480
08JAN96	AUSTIN	1980	1460	2085	1500	2350	2100	2025
	SAN ANTONIO	25200	650	5400	29000	21500	50000	44000
15JAN96	AUSTIN	2285	1950	1900	2400	1435	1645	3545
	SAN ANTONIO _	11400	50000	6500	19000	5220	810	3200
22JAN96	AUSTIN	875	1520	1655	425	330	600	180
	SAN ANTONIO	360	8000	4220	200	150	500	370
29JAN96	AUSTIN	45	40	25	80	20	20	40
	SAN ANTONIO	260	100	970	160	0	50	0
05FEB96	AUSTIN	135	25	260	45	20	25	18
	SAN ANTONIO	0	1220	1570	250	200	0	0
12FEB96	AUSTIN	18	0	0	Ð	0	0	0
	SAN ANTONIO	0	0	0	60	320	Ŏ	ŏ

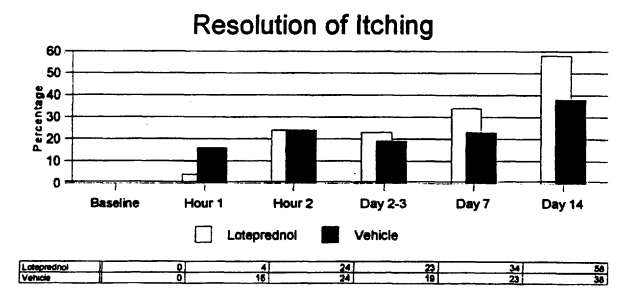
AGE		LOTEPREDNOL	PLACEBO	INV 176	INV 179	INV 180	INV 181
	N MEAN SD MIN MAX	67 39.2 10.5	68 38.2 9.3	37 39.5 11.0	36 39.2 7.9	32 37.2 11.1	30 38.8 9.7
GENDER							
MALE FEMALE	- N I	36 542	31 46% 37 54% p=1.000	19 511	15 421 21 581 QUAL p=0.8	16 501 16 501 81	
RACE	_						
CAUCASIAN HISPANIC OTHER	N I N I N I	45 672 17 253 5 72 HO:LE=PL	41 60% 25 37% 2 3% p=0.224	23 621 12 321 2 51 HO: INV E	15 42% 18 50% 3 8% QUAL p=0.04	24 751 7 221 1 31	5 172
IRIS							
L IGHT DARK	N I	31 46% 36 54% HO:LE-PL	28 413 40 593 p=0.605	17 46% 20 54% HO: INV E	9 25% 27 75% QUAL p=0.03	19 59% 13 41%	14 47% 16 53%
CITY							
AUSTIN SAN ANTONIO	N 1 N 1	15 22% 52 78% HO:LE=PL	15 22% 53 78% p=1.000		0 01 36 1001 WAL p=0.00	0 01 32 1001	30 1002 0 02
BASELINE POLLI	EN						
	N MEAN SD MIN MAX		68 119.5 010.3				30 1951.9 1058 9
5001-15000	N 2 N 3 N 2 N 2	8 12% 8 12% 17 26% 14 21% 19 29% HO:LE=PL p	10 15% .8 12% 17 25% 14 21% 19 28% 19 1762	7 19% 10 27% 0 0% 8 22% 12 32% HO: INV EQU	7 19% 1 3% 9 25% 13 36% 6 17% UAL p=0.000	0 01 4 132 0 01 7 232 20 652	4 133 1 32 25 832 0 02 0 02





A higher percentage of patients in the loteprednol group had resolution of redness compared to the vehicle group. The means scores are not impressively different.





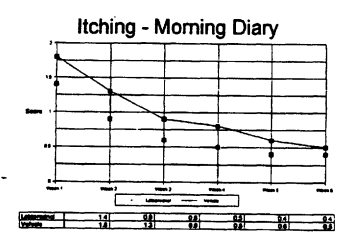
A significantly higher percentage of patients in the loteprednol group had resolution of itching compared to the vehicle group. The means scores are not impressively different.

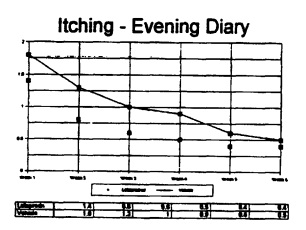
Bulbar Injection				OBSERVED RATINGS	. ME 14 05	EVEC ION.OF.		
and an injection	YI	SIT 1 (DAY D	1	VISIT 2	VISIT I	1153 (00-05). 11517 4	'4 VISIT 5	91517 6
	BASELINE	HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42
LOTEPREDNOL (LE)				S. 1-3		P	UA1 20	DA1 42
DISTRIBUTION	N 3	N 1	N 3	N 3	N 3	N 2	N 3	N 2
O:ABSENT	0 01	0 0%	2 33	14 22%	18 28%	23 364	6 264	2 403
0.5-1:MILD	0 04	19 283	33 492	34 521	40 623	32 SOR	15 653	3 601
1.5-2:HODERATE	48 72%	39 581	24 363	16 253	7 118	9 142	2 98	0 03
2.5-3:SEVERE	19 28%	9 132	8 12%	1 2%	0 03	0 01	0 03	0 0x
	67	67	67	65	65	64	23	5
MEAN	2.2	1.7	1.4	1.0	0.8	0.7	0.7	0.4
STANDARD ERROR	nα	በነ	A 1	A 1	0 1	0.1	0.1	0.5
MIN. MAX								
MEDIAN	4. U	2.U	1.0	1.0	1.0	1.0	1.0	U.5
PLACEBO (PL)								
DISTRIBUTION	N S	N 2	N %	H 2	H %	N \$	H \$	H \$
Q:ABSENT	0 01	0 0%	0 03	6 91	4 62	10 15%	2 103	0 03
0.5-1:MILD	0 0%	18 263	20 29%	32 481	33 50%	30 452	10 50%	31002
1.5-2:HODERATE	51 758	40 59%	36 53%	26 391	21 328	22 333	5 40%	0 62
2.5-3:SEVERE	17 254	10 15%	12 183	3 48	8 12%	4 63	0 01	0 02
N	68	68	68	67	66	66	20	3
HEAN	2.2	1.8	1.8	1.3	1.4	1.2	1.2	1.0
STANDARD ERROR	D. 0	0.1	0 1	A 1	N 1	.0.1	6.1	0.0
HIN. HAX								
MEDIAN	4.4	- 4.0	4.U	1.0	1.0	1.0	1.0	1.0
		VISIT 1	CHAI	GE FROM BASELING VISIT 2				
LOTEPREDNOL (LE)		HOUR 1	HOUR 2		VISIT 3 DAY 7	VISIT 4	VISIT 5	VISIT 6
FREQUENCY DISTRIBUTION		N S	N S	DAY 2-3		DAY 14	DAY 28	DAY 42
IMPROVED -3		0 03	0 03	2 31	N %	N 1	H \$	H \$
-2.5 · -2		0 03	4 63			5 82	2 9%	0 03
-1.51		27 40%	38 57%	20 311	22 34%	23 361	9 391	3 601
UNCHANCED -0.5 - 0.5		40 60%		31 481	36 558	30 47%	11 483	2 40%
1.5 - 1		0 02	25 37%	12 183	4 62	6 92	1 43	0 0%
MORSENED 2.5 2		0 01	0 0%	0 0% 0 0%	0 03	0 0% 0 0%	0 01	0 03
N	67	67	67				_0 OX	0 03
MEAN	2.2	-0.5	-0.8	65 -1.3	65	64	ສຸ	5
STANDARD ERROR	0.0	0.1	0.1	-1.3 ft 1	-1.5 0.1	-1.5 n 1	-1.7 n 2	-1.8 0.3
MIN. MAX	0.0	0.1	U. I	** 1	" '	21 1	пэ	n 4
HEDIAN	4.4	·u.a						
		-0.5	-1.0	-1.0	-1.5	-1.5	-1.5	-2.0
PLACEBO (PL)								
FREQUENCY DISTRIBUTION		N 2	N 3	N 3	N X	N X	N 3	N 3
IMPROVED -3		0 03:	0 03	1 13	1 23	1 23	0 03	0 0%
-2.52		0 0%	0 0%	8 123	4 68	14 213	2 103	0 03
-1.51		22 32%	23 343	34 513	39 59%	28 423	14 70%	31003
UNCHANGED -0.5 - 0.5		46 663	45 662	24 36%	20 303	22 333	4 203	0 03
1.5 - 1		0 02	0 0%	0 04	2 38	1 2%	0 03	0.01
MORSENED 2.5 - 2		0 0%	0 04	0 03	0 03	0 01	0 03	0 0%
K	68	66	68	67	66	66	20	3
HEAN	2.2	-0.4	-0.4	-0.9	-0.8	-1.0	-1.1	-1.0
STANDARD ERROR	0.0	0.1	0.1	01	D 1	D 1	0.1	0.0
MIN, MAX								
MEDIAN	4.∪	U.U	U.U	-1.U	-1.0	-1.U	-1.U	-1.0
TREATMENT EFFECT		-0.29		-0.				
958 CONF LIMITS		-0.43, -0.1	4	-0.67	0.38			
INVEST p-VALUE(c)	0.001							
TRT p-VALUE[d]	0.822	0.D54	0.000	0.001	0.000	0.000	0.010	0.087
TREATMENT EFFECT(e)	0.0	0.0	-0,5	-0.5	-0.5	-0.5	-0.5	-1.0
	0.0, 0.0	0.0. 0.0	0.5, 0.0	·0.5. 0.0 -	1.00.5	-1.0. 0.0	-1.0. 0.0	-1.5. O.G
(a) OBSERVED RATINGS ARE	THE HEAN	OF BOTH EYES	S AT THE V	isit;change from	BASELINE	IS OBSER VED F	Matjing - Basei	LINE MATING:
THUS. IMPROVENENT IS	A NEGATIV	E NUMBER						

Visits 5 and 6 do not have sufficient numbers of patients for evaluations of efficacy.

•			neserved R	ATINGS: HEAN	OF EYES (OO	-05 1/2		
Iteming	¥1511	1 (DAY 0)	0000	¥1517 2	A1211 3	41211 4		YISI7 6
	BASELINE	HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42
LOTEPREDNOL (LE)					* 3	н 5	H 2	H 2
DISTRIBUTION	N 3	N \$	N 3	N \$	N B 22 341	37 583	14 618	1 20%
Q:ABSENT	0 03	3 4%	16 248	15 233 20 318	21 321	18 281	6 268	3 601
0.5-1:TRACE	0 02	19 28%	24 36%	23 353	17 262	8 133	3 138	1 20%
1.5-2:MILD	0 02	25 37%	20 30%	7 118	5 8%	1 23	0 0%	0 0%
2.5-3:HODERATE	0 01	18 271	4 63	0 03	0 03	0 03	0 0%	0 0%
3.5-4:SEVERE	67100%	2 3%	3 4% 67	66	66	64	23	5
×	67	67	1.2	1.3	1.0	0.6	0.5	1.0
HEAN	4.0 a.n	1.8	1.6 N 1	0.1	0.1	0 1	0.1	r פ
STANDARD ERROR	41 11	•••						
MIN. MAX	4.0	2.0	1.0	1.0	1.0	Ų. U	U.U	1.0
MEDIAN	4.0	2.0	•.•	•				
PLACEBO (PL)							H 2	H 1
DISTRIBUTION	N 1	N 2	N 3	H \$	N 1	N 1		2 67%
O: ABSENT	0 0%	11 163	16 243	13 198	15 233	25 381	11 55% 5 25%	1 333
0.5-1:TRACE	0 02	21 31%	23 34%	18 27%	17 263	19 294	3 153	0 03
1.5-2:MILD	0 0%	21 31%	20 29%	19 283	20 303	17 26% 4 6%	1 53	0 03
2.5-3:MODERATE	0 0%	11 164	8 123	14 21%	12 1 83 2 31	1 23	0 03	0 03
3.5-4: SEVERE	68100%	4 68	1 1%	3 41		66	20	3
×	66	68	66	6)	66 1.5	1.0	0.7	0.3
HEAN	4.0	1.6	1.3	1.6	0.1	7.1	0.0	0.0
STANDARD ERROR	0.0 -	0 1	N 1	•••				
MIN. MAX				2.0	1.8	1.0	0.0	0.0
HEDIAN	4.0	1.5	. 1.0	2.0				
			CHAN	GE FROM BASE	LINE (OBSERV	ED - BASELIN	E (a))	
		VISIT 1		VIS17 2	VIS1T 3	VISIT 4	VISIT 5	V151T 6
LOTEPREDHOL (LE)		HOUR 1	HOUR 2	DAY 2-3	DAY 7	DAY 14	DAY 28	DAY 42
FREQUENCY DISTRIBUTION	N	N I	N X	N Z	N 3	N X	N 2	N I
IMPROVED -4	.•	3 42	16 24%	15 233	22 34%	37 564	14 613	1 201
-3.53		19 263	24 36%	20 31%	21 323	18 283	6 26%	3 60t
-2.52		25 37%	20 30%	23 354	17 263	8 132	3 134	1 20%
-1.51		18 27%	4 63	7 118	5 81	1 23	0 04	0 0%
UNCHANGED +0.5 - 0.5		2 3%	3 4%	0 04	0 0%	0 0%	0 0%	5
N	67	67	67	65	66	64	23	-3.0
MEAN	4.0	-2.2	-2.8	-2.7	-3.0	-3.4	-3.5 0.1	-3.U
STANDARD ERROR	0.0	n 1	0 1	n 1	6 1	n 1	•••	
MIN. MAX							-4.0	-3.0
MEDIAN	4.D	-2.0	-3.0	-3.0	-J. U	-4.U	-4.0	-0.0
PLACEBO (PL)		. 1	N 3	W 3	W 2	H 3	N 3	N 2
FREQUENCY DISTRIBUTION		11 16%	16 243	13 193	15 23%	25 363	11 55%	2 67%
IMPROVED -4		21 313	23 348	18 27%	17 268	19 294	5 25%	1 331
-3.53		21 313	20 29%	19 283	20 303	17 264	3 15%	0 03
-2.5 · ·2		11 163	8 123	14 21%	12 183	4 65	. 1 5%	0 0%
-1.51 UNCHANGED -0.5 - 0.5	:	4 63	1 18	3 48	2 33	1 23	0 03	0 0%
N	66	66	66	67	66	66	20	3
HEAN	4.0	-2.4	-2.7	-2.4	-2.5	-3.0	-3.3	-3.7 n a
STANDARD ERROR	0.0	0.1	0 1	n 1	n 1	n 1	0.3	., 4
MIN. MAX							4.0	.4.6
HEDIAN	4.0	٠٤.٦	٠.٥.٥	-6.U	-2.5	-3.0	-4.0	-4.0
TREATMENT EFFECT		0.09			0.40			
958 CONF LINITS		-0.23, 0.			90.11	west 4	VISIT 6	VISIT 6
UNIVARIATE AWALYSES		HOUR 1	HOUR 2	VISIT		0.004	0.799	0.157
TRT p-VALUE(c)		0.072	0.527	0.140	0.014	0.0	0.0	1.0
THEATHENT EFFECTED)		0.0	0.0	0.0	-0.5	-1.0. 0.0	0.0, 0.0	0.0, 2.0
952 CONF LIMITS		0.0. 0.5	0.0. 0.0	*1.0. U.V	INC THRUIMS	THE GENERE !	4) FOR ALL PA	
(a) OBSERVED RATINGS	ARE THE NE	WED BUTH !	Y SHI IA CST:	1511. BESEL IMPROPRIEMENT	IS A MEGATIN	E MANGER	.,	
(a) DESERVED HATTHIS CHANGE FROM BASEL	INE 15 DUSE	TOED HATING	· 4; IMA	THE MOST PLANT				

Visits 5 and 6 do not have sufficient numbers of patients for evaluations of efficacy.





The graphs show overall improvement from baseline in both groups and little difference between groups.

Patients with zero rating (sign or symptom no longer present)

		(agn or symptom no longer present							
			. \	/isit 1	Visit 2	Visit 3	Visit 4		
Measure:	Treatment	N	Hour 1	Hour 2	Day 2/3	Day 7	Day 14		
Discomfort:	LE	65	7%	21%	30%	48%	63%		
	Placebo	66	13%	24%	24%	32%	48%		
Foreign body sensation:	LE	61	24%	38%	54%	54%	72%		
	Placebo	63	30%	39%	43%	44%	53%		
Burning/Stinging:	LE	64	18%	32%	47%	56%	63%		
	Placebo	66	30%	28%	33%	42%	54%		
Photophobia:	LE	53	29%	40%	51%	57%	63%		
	Placebo	56	25%	36%	39%	40%	56%		
Tearing:	LE	63	20%	42%	56%	60%	81%		
	Placebo	62	30%	37%	50%	49%	66%		
Discharge:	LE	31	45%	52%	68%	71%	81%		
	Placebo	24	44%	32%	50%	48%	67%		
Palpebral injection:	LE	64	2%	3%	5%	16%	21%		
	Placebo	67	1%	1%	1%	3%	9%		
Chemosis:	LE	53	0%	4%	23%	25%	35%		
	Placebo	60	0%	2%	22%	27%	41%		
Erythema:	LE	50	13%	21%	40%	54%	55%		
	Placebo	53	4%	11%	28%	28%	37%		

N is the intent-to-treat sample size of patients with the sign or symptom present at baseline.

Reviewer's Comments:

With the exception of the 1st visit and Chemosis, the loteprednol group had higher percentages of symptom resolution.

Intraocular Pressure:

		Number of patients					
Treatment	Elevation in IOP (mm Hg)	Day 7	Day 14	Day 28	Day 42		
LE	> 15	0	0	0	1		
	10-15	0	0	0	0		
	6-9	3 1 4 62 63 60	4				
	< 6	62	63	60	59		
Placebo	> 15	0	1	0	o		
	10-15	0 0 0 0 0 0 3 1 4 62 63 60	0	0			
	6-9	0	0	0	0		
	< 6		65	64			

The distributions (above and below) displayed are the change in IOP in the eye with the greatest increase from baseline in IOP.

LOTEPREDNOL	BASELINE	VISIT 3 Day 7	VISIT 4 DAY 14		VISIT 6 Day 42	
DISTRIBUTION	N 1	N 2	N 2	N I	N 1	
< 20 MM HG	64 961	61 942	62 971	61 951	63 981	
20 - 25 MM HG	3 41	4 62	2 38	3 51	0 01	
26 - 31 MM HG	0 01	0 01	0 01	0 01	0 01	
> 31 MM HG	0 02	0 02	0 02	0 01	1 21	
N	67	65	64	64	63	
MEAN	14.9	15.5	15.3	16.0	15.6	
STANDARD ERROR	. 0.3	0.3	0.3	0.3	0.4	
MIN. MAX			•••		•••	
MEDIAN	15.0	15.0	15.0	16.0	15.0	
PLACEBO (PL)						
< 20 MM HG	67 991	661001	59 891	62 95%	63 98%	
20 - 25 MH HG	1 12	8 01	6 92	3 5%	1 21	
26 - 31 MM HG	0 02	0 02	0 02	0 02	0 01	
> 31 MM HG	0 01	0 01	1 23	0 02	0 01	
N	68	66	66	65	64	
MEAN	15.7	15.1	15.6	15.2	14.5	
STANDARD ERROR HJN. MAX	0.3	0.3	0.4	0.3	0.3	
MEDIAN	16.0	15.5	15.3	15.0	14.5	

Reviewer's Comments:

Elevation in IOP was seen more frequently in the loteprednol group.

Adverse Experiences: (Greater than 2%)

SPECIAL SENSES	- PATIENTS		IENTS ING EVENT	TOTAL Number	SEVERITY OF EVENTS		
,	AT RISK		AST ONCE	OF EVENTS	HILD	MODERATE	SEVERE
BODY AS A WHOLE -Any event			•				
LOTEPREDNOL	67	18	272	27		10	_
PLACEBO	68	15	222	27 19	13 13	10 5	4
	-	10	764	17	13	•	1
RHINITIS (NOSE)							
LOTEPREDNOL	67	16	243	19	9	9	1
PLACEBO	68	9	131	12	4	6	2
ITCHING. EYE (EYE/GEN)							
LOTEPREDNOL	67	11	150	••	_	_	_
PLACEBO	68	11	163	13	7	5	1
FERGEDO	90	8	12\$	9	6	2	1
HEADACHE (HEAD)							
LOTEPREDNOL	67	10	15%	15	9	4	2
PLACEBO .	68	8	12%	11	ź	3	ī
CUPAGE IF AFAF (CON.)						-	-
CHEMOSIS (EYE/CONJ)							
. LOTEPREDNOL	67	9	132	9	8	0	1
PLACEBO	68	11	161	13	10	2	1
DISCHARGE. EYE (EYE/GEN)							
LOTEPREDNOL	67	8	12%	9	4	•	•
PLACEBO	68	9	13%	9	8	3	0
. 510550	V.	7	134	,	0	1	0
BURNING/STINGING, EYE, NO		ATION (EYE/GEN)				
LOTEPREDNOL	67	6	91	7	4	2	1
PLACEBO	68	5	72	5	3	ĩ	1
EPIPHORA (EYE/APP)							
LOTEPREDNOL	67	_		_			
	67	5	71	5	4	0	1
PLACEBO	68	9	131	12	7	5	Q
COUGH INCREASED (GEN)							
LOTEPREDHOL	6 7	4	61	5	2	3	0
PLACEBO	68	2	31	2	ī	ĭ	ŏ
EYE/VISION. BLURRED (EYE	/VIS)						
LOTEPREDNOL	67	4	61	7	6	1	0
PLACEBO	68	2	31	Ž	ì	i	Ö
DISCOMFORT, EYE (EYE/GEN)							
LOTEPREDNOL	67	•	10	•	_	_	_
PLACEBO		3	43	3	1	2	0
PORCEBU	68	2	31	2	2	0	0
INFECTION (GEN)							
LOTEPREDNOL	. 67	3	41	4	2	2	0
PLACEBO	68	Ö	02	Ö	Õ	õ	Ö
		-		•	•	•	•
FOREIGN BODY SENSATION (EY LOTEPREDNOL		•	45	_		_	
	67	3	413	3	1	2	0
PLACEBO	68	4	61	4	2	1	1

INJECTION (EYE/CON)							
LOTEPREDNOL	67	3	42	3	3	0	0
PLACE80	68	15	221	19	16	3	0
PHOTOPHOBIA (EYE/VIS)							
LOTEPREDNOL	67	3	42	3	0	3	0
PLACEBO	68	1	12	1	0	1	0
PHARYNGITIS (NASP)							
LOTEPREDNOL	67	3	42	3	2	1	0
PLACEBO	68	2	31	2	1	1	0
ACCIDENTAL INJURY (GEN)							
LOTEPREDNOL	67	2	31	2	1	0	1
PLACEBO	68	0	02	0	0	0	0
FACE EDEMA (HEAD)							
LOTEPREDNOL	67	2	32	2	0	1	1
PLACEBO	68	2	31	2.	1	1	0
DIARRHEA (EC)							
LOTEPREDNOL	67	2	31	2	0	2	0
PLACEBO	68	0	02	0	0	0	0
VOMITING (GEN)							
LOTEPREDNOL	67	2	3\$	2	1	1	0
PLACEBO .	68	0	02	0	G	0	0
ASTHMA (BRON)							
LOTEPREDNOL	67	2	31	2	0	2	0
PLACEBO	68	0	OZ	0	0	0	0
SINUSITIS (SINS)							
LOTEPREDHOL	67	2	32	2	0	2	0
PLACEBO	68	1	12	1	0	1	0

Reviewer's Summary of Safety and Efficacy

Marginal efficacy has been demonstrated in the resolution of itching and redness. Adverse experiences in this limited study (42 days) were generally confined to mild to moderate ocular events. There was an increased chance of increased IOP during use.

8.1.3 Study #3

Protocol #141

Title:

Efficacy and Safety of Lotemax™ BID vs Lotemax™ QID in the

Antigen Challenge Model of Acute Allergic Conjunctivitis

Investigators:

Mark Abelson, M.D. (Investigator #108)
ORA Clinical Research and Development

863 Tumpike Street

North Andover, MA 01845

Objective:

To compare two dose regimens of loteprednol etabonate 0.5% ophthalmic suspension on the prevention of signs and symptoms induced by an ocular antigen challenge, and to evaluate the

duration of action of this effect.

Study Design:

A randomized, double-masked, placebo controlled, paired

comparison, single center study.

Population:

There were 60 otherwise normal adults with known allergies to

specific antigens.

Schedule:

All subjects received loteprednol etabonate 0.5% ophthalmic suspension in one eye and vehicle placebo in the contralateral eye. Study drugs were instilled either BID or QID for 28 days, from Day 7 to Day 35. Visits and antigen challenges were carried out

on Days 0, 7 (baseline), 21 and 35.

Study Plan

Study 141, was a prospective, double masked, placebo controlled, single center, paired-comparison of loteprednol etabonate 0.5% ophthalmic suspension (BID or QID) versus placebo (vehicle) in the antigen challenge model of acute allergic conjunctivitis. Sixty (60) subjects who had a minimum pre-determined response to an ocular antigen challenge were enrolled in the study. All subjects received drug in one eye and vehicle in the contralateral eye. Subjects were randomized with respect to which eye received active drug. The first 30 patients received treatment in a BID dosing schedule and the second 30 patients were on a QID dosing schedule.

On Day 0, a conjunctival allergen test was performed bilaterally using allergen to which the subject had a history of sensitivity (weed, animal dander, tree or grass) diluted with phosphate buffered saline. Doses ranging from 19 to 1250 allergen units per 25 µL dose were administered in a dose related manner until a response of 2+ itching and redness at 10 minutes post instillation was achieved. If the maximum dose was reached without achieving this response the subject was excluded from further study participation and an exit form was completed. Subjects who tested positively were asked to return for the Visit 2 qualifying challenge. On Day 7, the subjects were challenged with the highest dose of allergen used on Day 0 to ensure that their response was still present. Subjects who qualified by their response to the second challenge were to begin a twenty eight (28) day period of study medication use. Loteprednol etabonate 0.5% ophthalmic suspension and vehicle placebo were to be instilled into the appropriate eye according to either a BID or QID dosing schedule. Subjects were rechallenged on Day 21, at 15 minutes after the latest dose of test article and on Day 35, subjects were randomly divided into two groups for challenges at 2 hr or 8 hr after the final dose of test article.

Ocular safety evaluations included an external examination, slit lamp examination, tonometry and visual acuity taken prior to enrollment and at scheduled times during the study. Systemic safety evaluations were treated by subject comment with physician follow-up.

Inclusion Criteria

- 3. 18 years of age or older, of either sex and of any race.
- 4. Manifest a successful challenge, inducing at least 2+ itching and 2+ redness bilaterally.
- A positive history of allergy to grasses, animal dander, weeds, or trees. Positive skin tests, prior positive reactions to allergen challenge or verbal subject report consistent with allergy will constitute a positive history.

Exclusion Criteria

- 1. Contraindications to the use of the study medication(s).
- 2. Known sensitivity or allergy to the study drug(s) or their components.
- 3. Presence of any significant illness that could be expected to interfere with the study, particularly any autoimmune disease, e.g., rheumatoid arthritis.
- 4. Presence of bacterial or viral ocular infection.
- 5. Presence of blepharitis, follicular conjunctivitis, iritis, or preauricular lymphadenopathy.
- Presence of mucous discharge, excess lacrimation, or burning as symptoms of ocular disease (possible dry eye).
- 7. History of dry eye or evidence of dry eye demonstrated by slit lamp examination.
- 8. Manifest signs and symptoms of clinically active allergic conjunctivitis (>1+ redness and/or the presence of any itching) at the baseline eye examination at visits 1 and 2.

- Use of ocular medications of any kind, including tear substitutes, or systemic medication that may interfere with a normal vasodilatory response or with normal lacrimation for an appropriate wash-out period prior to the start of the study and for the duration of the study (i.e., non-steroidal anti-inflammatories, anti- histamines, etc. within 72 hours, corticosteroids within 7 days, mast cell stabilizers within 14 days).
- 10. Contact lenses worn 3 days prior to or during study period.
- 11. Pregnant or nursing women; or women of childbearing potential who test positive to a pregnancy test.
- 12. Participation in a clinical trial or use of an investigational drug or device within the last 30 days.

Efficacy Criteria

Intraocular differences in itching and mean redness (the mean score of redness in the ciliary, episcleral and conjunctival vessel beds) were the primary efficacy variables. Secondary variables included chemosis, tearing, lid swelling and mucous discharge.

Most signs and symptoms were rated on a four point scale (0-4) where 0= absent, 1= mild, 2= moderate, 3= severe and 4= unusually severe. Increments of 0.5 units were also assessed, e.g., a score of 1.5 would rate between mild and moderate. Tearing was rated on a 0-3 scale where 0= none, 1= mild (eyes felt slightly watery), 2= moderate (blows nose occasionally) and 3= severe (tears rolling down cheeks).

For itching expanded definitions were provided

0 ≈ none

1 = an intermittent tickle sensation in the inner corner

2 = a mild continuous itch, not requiring rubbing

3 = a definite itch; you would like to be able to rub

4 = an incapacitating itch which would require eye rubbing

Demographics

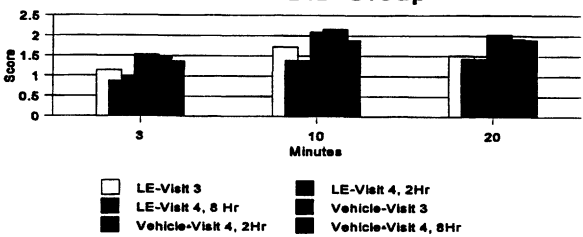
Mean Age 33.2 ± 10.8 years (min=19, max=85)

 Gender
 Male=33 (55%)
 Female=27 (45%)

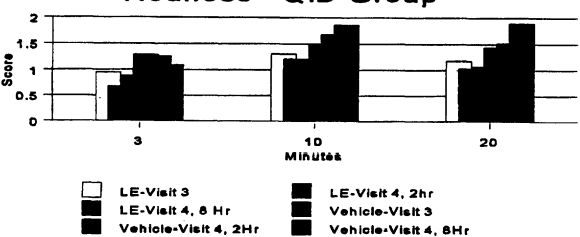
 Race
 Caucasian=58 (97%)
 Hispanic=2 (3%)

 Iris Pigmentation
 Light=36 (60%)
 Dark=24 (40%)



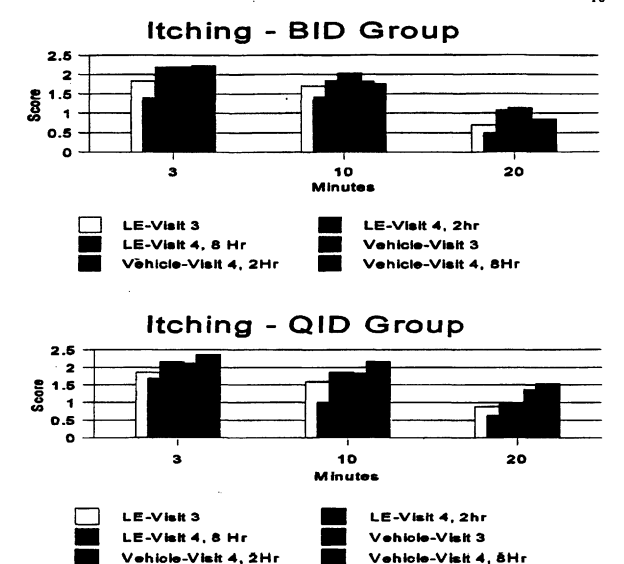


Redness - QID Group



Reviewer's Comments:

There are minimal differences between groups, although the loteprednol eye generally does better than the vehicle eye.



There are minimal differences between groups, although the loteprednol eye generally does better than the vehicle eye.

Visit	3 -B	ID G	roup
--------------	------	------	------

Visit 3 -BID Group					C-65:	.	Division
	LE Ey (n = 2 Mean	28)	Vehic (n = 2 Mean	•	Effica Score (n = 2 Mean	8)	P-Value (2-Tail)
Mean Redness							
3 Min Post Challenge	1.14	0.81	1.53	0.82	-0.39	0.71	0.0077
10 Min Post Challenge	1.72	0.91	2.10	0.74	-0.38	0.91	0.0381
20 Min Post Challenge	1.51	1.00	2.03	0.84	-0.52	1.04	0.0128
ttching							
3 Min Post Challenge	1.84	1.16	2.20	0.83	-0.36	0.96	0.0596
10 Min Post Challenge	1.70	1.00	2.04	0.83	-0.34	0.89	0.0544
20 Min Post Challenge	0.70	0.72	1.14	0.61	-0.45	0.63	0.0008
Visit 3 -QID Group							
Mean Redness							
3 Min Post Challenge	0.95	0.75	1.29	0.88	-0.34	0.56	0.0035
10 Min Post Challenge	1.31	0.95	1.50	0.94	-0.19	0.84	0.2385

Wadii Vanices								
3 Min Post Challenge	0.95	0.75	1.29	0.88	-0.34	0.56	0.0035	
10 Min Post Challenge	1.31	0.95	1.50	0.94	-0.19	0.84	0.2385	
20 Min Post Challenge	1.18	1.01	1.44	1.06	-0.26	0.80	0.0928	
Itching					•			
3 Min Post Challenge	1.86	0.94	2.12	0.99	-0.27	0.90	0.1259	
10 Min Post Challenge	1.59	0.92	1.84	0.95	-0.25	0.81	0.1144	
20 Min Post Challenge	0.89	0.99	0.98	1.04	-0.09	0.84	0.5782	

^{**} Efficacy Score = (Difference Score - Treated vs Vehicle Eye)

	LE Ey Mean		Vehic Mean	le Eye Sd	Effica Score Mean	**	P-Value (2-Tail)
N=15							
Mean Redness							
3 Min Post Challenge	0.88	0.72	1.47	0.97	-0.59	1.07	0.0510
10 Min Post Challenge	1.39	0.79	2.17	0.78	-0.78	0.95	0.0070
20 Min Post Challenge	1.43	0.90	1.93	1.04	-0.50	1.15	0.1132
Itching		•					
3 Min Post Challenge	1.40	1.20	1.90	0.95	-0.50	1.07	0.0916
10 Min Post Challenge	1.37	0.93	1.83	0.77	-0.47	0.90	0.0632
20 Min Post Challenge	0.50	0.60	0.83	0.70	-0.33	0.75	0.1064

Visit 4 - BID Group, 8 Hour Challenge

N=13 Mean Redness							
3 Min Post Challenge	0. 9 6	0.87	1.37	0.96	-0.41	0.72	0.0620
10 Min Post Challenge	1.24	0.94	1.90	1.12	-0.65	0.81	0.0133
20 Min Post Challenge	1.29	1.11	1.91	1.17	-0.62	0.86	0.0236
Itching							
3 Min Post Challenge	2.19	0.97	2.23	0.93	-0.04	0.25	0.5845
10 Min Post Challenge	1.85	0.69	1.77	0.88	0.08	0.93	0.7711
20 Min Post Challenge	1.08	1.12	0.85	0.66	0.23	1.15	0.4824

Visit 4 - QID	Group,	2 Hour	Challenge
---------------	--------	--------	-----------

	LE Ey Mean		Vehici Mean	•	Effica Score Mean	**	P-Value (2-Tail)
N=14 Mean Redness 3 Min Post Challenge 10 Min Post Challenge 20 Min Post Challenge	0.68	0.63	1.26	0.85	-0.58	0.55	0.0017
	1.21	0.83	1.68	1.04	-0.46	0.81	0.0512
	1.04	0.92	1.52	0.98	-0.49	0.73	0.0264
ttching 3 Min Post Challenge 10 Min Post Challenge 20 Min Post Challenge	1.68	1.15	1.75	1.20	-0.07	0.55	0.6349
	1.00	0.88	1.61	1.10	-0.61	1.26	0.0 9 43
	0.64	0.60	1.36	0.95	-0.71	0.97	0.0168

Visit 4 - QID Group, 8 Hour Challenge

• •			-		Effica	CV	P-Value
	LE Ey			le Eye	Score	**	(2-Tail)
	Mean	50	Mean	Sa	Mean	Sa	
N=12							
Mean Redness							
3 Min Post Challenge	0.88	0.79	1.10	0.69	-0.22	0.71	0.3004
10 Min Post Challenge	1.19	0.97	1.86	0.67	-0.67	0.75	0.0104
20 Min Post Challenge	1.07	1.04	1.90	0.83	-0.83	0.75	0.0027
Itching							
3 Min Post Challenge	2.17	0.94	2.37	0.91	-0.21	0.40	0.0960
10 Min Post Challenge	1.87	1.09	2.17	1.21	-0.29	0.54	0.0891
20 Min Post Challenge	0.96	0.89	1.54	1.20	-0.58	1.02	0.0728

Mean Intraocular Pressure (mmHg)

Visit	Day	N	LE Eye Mean SD	Range	Vehicle Eye Mean SD	Range
BID Dosing	Group					
2	7	28	15.04 2.69	10.0 - 21.0	14.93 2.88	10.0 - 21.0
3	21	28	14.43 2. 9 0	8.0 - 22.0	14.25 2.82	7.0 - 20.0
4 (2hr)	35	15	15.93 2.76	12.0 - 20.0	15.93 2.66	12.0 - 20.0
4 (8hr)	35	13	14.54 2.63	11.0 - 21.0	14.85 2.48	11.0 - 20.0
QID Dosing	Group	,				
2	7	29	15.38 2.47	11.0 - 18.0	15.38 2.51	11.0 - 18.0
3	21	29	14.38 2.11	11.0 - 16.0	14.41 2.56	11.0 - 20.0
4 (2hr)	35	14	14.93 2.87	11.0 - 17.0	14.43 2.68	10.0 - 17.0
4 (8hr)	35	12	15.00 2.86	10.0 - 19.0	14.83 2.89	10.0 - 18.0

Reviewer's Comments:

There were no elevations above 10 mmHg in either group.

Adverse Events:

Headaches were the most commonly reported events during this study.

Reviewer's Summary of Safety and Efficacy

Marginal efficacy has been demonstrated in the relief of itching and redness. Adverse experiences cannot be well determined from this study.

8.1.4 Study #4 Protocol # 145

Title: Comparison of dose regimen study assessing the efficacy of

various concentrations of loteprednol etabonate ophthalmic suspension in the antigen challenge model of acute allergic

conjunctivitis.

Objective: Paired Comparison Study: To compare three doses of

loteprednol etabonate (0.1%, 0.2% and 0.3%) ophthalmic suspension on the prevention of signs and symptoms

induced by an ocular antigen challenge, and to evaluate the

duration of action of this effect.

Parallel Group Study: To evaluate the loteprednol

etabonate 0.5% ophthalmic suspension compared to vehicle placebo on the prevention of signs and symptoms induced

by an ocular antigen challenge.

Study Design: Study 145, was a prospective, double masked, placebo

controlled, single center, study which consisted of two

separate parts.

Paired Comparison Study: A randomized, double-masked,

placebo controlled, paired comparison, single center study

(0.1%, 0.2% and 0.3%).

Parallel Group Study: A randomized, double-masked,

placebo controlled, parallel group, single center study

(0.5%).

Population: There were 120 otherwise normal adults with known

allergies to specific antigens.

Test Drug Schedule:

Paired Comparison Study: All subjects received loteprednol etabonate ophthalmic suspension (either 0.1%, 0.2% and 0.3%) in one eye and placebo (vehicle) in the contralateral eye. Subjects were randomized with respect to which eye received which drug.

Parallel group Study: Subjects received either active drug (0.5%) or placebo (vehicle) in both eyes.

For all subjects, study drugs were instilled QID for 28 days, from Day 7 to Day 35. Visits and antigen challenges were carried out on Days 0, 7 (baseline), 21 and 35.

Investigator

Mark B. Abelson, M.D.
ORA Clinical Research and Development
863 Tumpike Street
North Andover, MA 01845

Study Plan

One hundred and twenty (120) subjects who had a minimum pre-determined response to an ocular antigen challenge were enrolled in the study. Ninety (90) subjects were to receive drug in one eye and vehicle in the contralateral eye. These subjects were randomized with respect to which eye received active drug and to which dose of drug was received. Thirty (30) subjects were randomized to receive either 0.5% loteprednol etabonate ophthalmic suspension or vehicle placebo bilaterally.

All subjects were administered a pre-study challenge on Day 0 to determine their response to rising doses of allergen. On Day 7 the subjects were challenged with the highest dose of allergen used on Day 0 to ensure that their response was still present. Subjects who qualified by their response to the second challenge were to begin a twenty eight (28) day period of study medication use. In ninety (90) randomized subjects loteprednol etabonate ophthalmic suspension (0.1%, 0.2% or 0.3%) and placebo (vehicle) were to be instilled into the appropriate eye according to a QID dosing schedule. Subjects were rechallenged on Day 21 (14 days of treatment) at 30 minutes after the latest dose of test article and on Day 35, subjects were divided into two groups from a predetermined randomization for challenges at 2 hr or 4 hr after the final dose of test article. The remaining 30 subjects were randomized to receive either loteprednol etabonate 0.5% ophthalmic suspension or placebo (vehicle) bilaterally on a QID schedule. Subjects were rechallenged on Day 21 (14 days of treatment) at 30 minutes after the latest dose of test article and on Day 35 (28

days of treatment) at 2 hours after the final dose of test article.

Ocular safety evaluations included an external examination, slit lamp examination, tonometry and visual acuity taken prior to enrollment and at scheduled times during the study. Systemic safety evaluations were treated by subject comment with physician follow-up.

Inclusion Criteria

- 18 years of age or older.
- Of either sex and of any race.
- Manifest a successful challenge, inducing at least 2+ itching and 2+ redness bilaterally.
- A positive history of allergy to grasses, animal dander, weeds, or trees. Positive skin tests, prior
 positive reactions to allergen challenge or verbal subject report consistent with allergy will
 constitute a positive history.

Exclusion Criteria

- Contraindications to the use of the study medication(s).
- Known sensitivity or allergy to the study drug(s) or their components.
- Presence of any significant illness that could be expected to interfere with the study, particularly any autoimmune disease, e.g., meumatoid arthritis.
- Presence of bacterial or viral ocular infection.
- Presence of biepharitis, follicular conjunctivitis, iritis, or preauricular lymphadenopathy.
- Presence of mucous discharge, excess lacrimation, or burning as symptoms of ocular disease (possible dry eye).
- History of dry eye or evidence of dry eye demonstrated by slit lamp examination.
- Manifest signs and symptoms of clinically active allergic conjunctivitis (>1+ redness and/or the presence of any itching) at the baseline eye examination at visits 1 and 2.
- Use of ocular medications of any kind, including tear substitutes, or systemic medication that
 may interfere with a normal vasodilatory response or with normal lacrimation for an appropriate
 wash-out period prior to the start of the study and for the duration of the study (i.e., non-steroidal
 anti-inflammatories, anti- histamines, etc. within 72 hours, corticosteroids within 7 days, mast cell
 stabilizers within 14 days).
- Contact lenses worn 3 days prior to or during study period.
- Pregnant or nursing women; or women of childbearing potential who test positive to a pregnancy test.
- Participation in a clinical trial or use of an investigational drug or device within the last 30 days.

Efficacy Criteria

For subjects in the paired comparison arm of the study interocular differences (efficacy scores) in itching and mean redness (the mean score of redness in the ciliary, episcleral and conjunctival vessel beds) were the primary efficacy variables. Secondary variables were interocular differences (efficacy scores) in chemosis, tearing and lid swelling. Mucous discharge was evaluated in all subjects. For the subjects in the parallel group comparison the same parameters were recorded, however the variables were the difference in mean scores between those subjects who received active medication and those who received placebo (vehicle).

Most signs and symptoms were rated on a four point scale (0 - 4) where 0 = absent, 1 = mild, 2 = moderate, 3 = severe and 4 = unusually severe. Increments of 0.5 units were also assessed, e.g., a score of 1.5 would rate between mild and moderate. Tearing was rated on a 0-3 scale where 0 = none, 1 = mild (eyes felt slightly watery), 2 = moderate (blows nose occasionally) and 3 = severe (tears rolling down cheeks). Mucous discharge was rated as either absent (0) or present (1).

For itching expanded definitions were provided

0 = none

1 = an intermittent tickle sensation in the inner comer

2 = a mild continuous itch, not requiring rubbing

3 = a definite itch; you would like to be able to rub

4 = an incapacitating itch which would require eye rubbing

Product Batches:

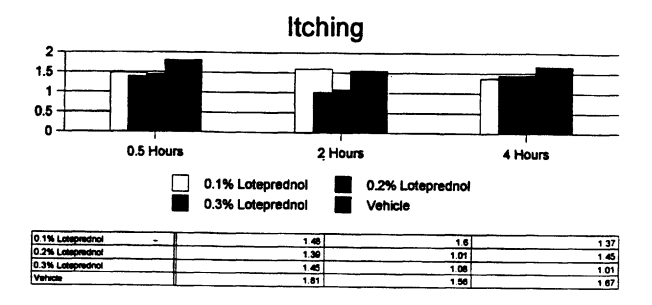
loteprednol etabonate 0.1% ophthalmic suspension - Batch # 004-95 - Batch # 002-95 loteprednol etabonate 0.3% ophthalmic suspension loteprednol etabonate 0.5% ophthalmic suspension - Batch # 001-95 - Batch # 001-93 - Batch # 001-93 placebo (vehicle) - for 0.5% - Batch # 002-93 - Batch # 003-95

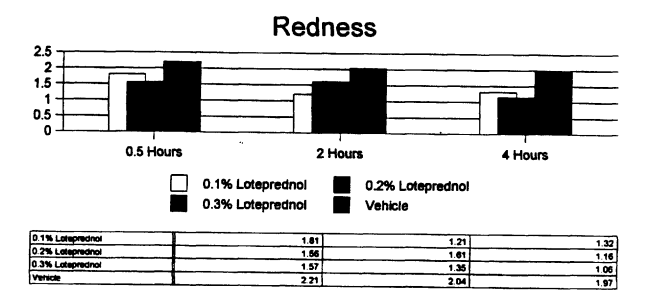
Procedure	Visit 1	Visit 2	Visit 3	Viola 4
l location	Day 0		1	Visit 4
•	Day U	Day 7	Day 21	Day 35
		(Day 7-9)	(Visit 2 + 14)	(Visit 3 + 14)
			(Day 21-23)	(Day 35-37)
Informed Consent	X	1		
inclusion/Exclusion	Х			
Demographics	Х			
Medical/Surgical History	X			
Medication History	Х	Х	X	Х
Urine Pregnancy Test	X			X
Ophthalmic Exam: Visual Acuity & Slit Lamp	X	X	X	X
Ocular Symptoms of Allergic Conjunctivitis	X	Х	Х*	X*
ntraocular Pressure		X	X	Х
Antigen Challenge	Х	Х	X	Х
Photograph **			X	Х
Assignment of subject to 2 hr or 4 hr challenge schedule				Х
Dispense Medication	•	Х		
Recover Medication				X
Exit Form				X

^{*} Evaluation of Allergic conjunctivitis at 3, 10 and 20 minutes after allergen challenge ** Photographs taken immediately after 10 minute evaluation

			Reas	son Study Incon	npiete
Treatment	Enrolled	Completed Study	Lack of Efficacy	Adverse Event	Other (Unrelated)
Paired Comparis	son				
Lotaprednoi etabonate 0.1% ophtheimic euspension	28	26 (93%)	0	0	2 (7%)
Loteprednol etabonate 0.2% ophthelmic suspension	31	31 (100%)	0	0	0
Loteprednoi etabonate 0.3% ophthelmic suspension	29	29 (100%)	0	0	0
Parallel Group					
Loteprednol etabonate 0.5% ophthelmic suspension	16	16 (100%)	0	0	0
Vehicle placebo	16	14 (88%)	0	0	2 (12%)

	All subjects		0.1% group	0.2% group	0.3% group	0.5% group
Age	n	120	28	31	29	32
(years)	mean	36.4±11	36.5±11	35.7±11	38.6±12	35.0±10
	Min-Max					
Gender						
Male	n (%)	61 (51%)	18 (64%)	13 (42%)	15 (52%)	15 (47%)
Female	n (%)	59 (49%)	10 (36%)	18 (58%)	14 (48%)	17 (53%)
Race	•					
Caucasian	n	119 (99%)	28 (100%)	31 (100%)	28 (97%)	32 (100%)
Hispanic	n	1 (1%)	0	0	1 (3%)	0
iris Pigmer	tation					
Light	n	80 (67%)	22 (79%)	20 (65%)	19 (66%)	19 (59%)
Dark	n	40 (33%)	6 (21%)	11 (35%)	10 (34%)	13 (41%)





All of the loteprednol groups perform slightly better than the vehicle group. The 0.3% loteprednol group is usually better than the other two groups although the differences are very small.

Itching	LE Eye	•	Vehick	e Eye	Efficac	y Sčôřě ™	p-vāluē
	Mean	SD.	Mean	Sd.	Mean	Sd.	(2-tail)
0.1% Group (n = 27)							
3 min post-challenge	1.93	0.97	2.19	0.94	-0.26	0.98	0.1828
10 min post-challenge	1.56	0.98	1.83	0.99	-0.28	1.13	0.2126
20 min post-challenge	0.96	0.89	1.17	1.13	-0.20	0.84	0.2162
0.2% Group (n = 31)							
3 min þóst-chállenge	1.79	1.32	2.27	1.06	-0.48	1.04	0.0144
10 min post-challenge	1.60	1.17	2.29	0.87	-0.69	0.92	0.0002
20 min post-challenge	0.79	0.92	1.24	1.02	- 0.45	1.01	0.0002
0.3% Group (n = 29)							
3 min post-challenge	2.07	1.00	2.40	0.95	-0.33	0.79	0.0321
10 min post-challenge	1.41	0.95	1.83	0.96	-0.41	0.78	0.0080
20 min post-challenge	0.88	0.91	1.09	1.01	-0.21	0.80	0.1728
Redness	155) fabrus	. =			
Neuliess	LE Eye	•	Vehicle	: Eye	ETTICAC	y Score**	p-value
	Mean	Sđ.	Mean	Sd.	Mean	Sd.	(2-tail)
0.1% Group (n - 27)							
3 min post-challenge	1.44	0.79	1.83	0.80	-0.40	0.74	0.0100
10 min post-challenge	2.00	0.86	2.20	0.90	-0.20	0.77	0.1961
20 min post-challenge	1.96	0.91	2.21	0.85	- 0.23	0.85	0.1636
0.2% Group (n - 31)							
3 min post-challenge	1.24	0.84	1.77	0.95	-0.54	0.63	1000.0
10 min post-challenge	1.73	1.11	2.37	0.96	-0.64	0.91	0.0005
20 min post-challenge	1.72	1.11	2.32	0.98	-0.60	0.87	0.0006
0.3% Group (n - 29)							
3 min post-challenge	1.20	0.85	1.99	0.93	-0.80	0.84	0.0001
10 min post-challenge	1.81	0.75	2.61	0.66	-0.80	0.68	0.0001
20 min post-challenge	1.70	0.91	2.59	0.75	- 0.89	0.86	0.0001

Rching- 2 Hr Challenge	LE Eye	,	Venicie	Eÿē	Efficac	y Score**	p-value
	Mean	Sd.	Mean	Sđ.	Mean	Sđ.	(2-tail)
0.1% Group (n = 12)							
3 min post-challenge	1.62	1.00	1.71	1.27	-0.08	0.58	0.6147
10 min post-challenge	1.92	1.08	1.96	1.16	-0.04	0.54	0.7949
20 min post-challenge	1.25	1.01	1.33	1.01	-0.08	0.47	0.5505
0.2% Group (n = 15)							
3 min post-challenge	1.63	1.17	2.07	0.78	-0.43	1.12	0.1548
10 min post-challenge	0.93	1.16	1.57	0.86	-0.63	1.17	0.0551
20 min post-challenge	0.47	0.74	0.63	0.84	-0.37	0.97	0.1662
0.3% Group (n = 15)							
3 min post-challenge	1.67	1.25	-2.13	1.08	-0.47	0.93	0.0737
10 min post-challenge	1.07	1.02	1.73	1.08	-0.67	1.01	0.0230
20 min post-challenge	0.50	0.73	0.73	0.86	-0.23	0.56	0.1306
Redness- 2 Hr Challenge	LE Eye		Vehick	a Fva	EMAAA	y Score**	p-value
			00111011	- - , ·		, 00010	(2-tail)
	Mean	Sd.	Mean	Sd.	Mean	Sd.	(2-141)
0.1% Group (n = 12)							
3 min post-challenge	0.72	0.71	1.25	0.91	-0.53	0.51	0.0044
10 min post-challenge	1.62	0.75	1.72	0.94	-0.10	0.76	0.6677
20 min post-challenge	1.29	0.92	2.06	0.93	• 0.76	0.84	0.0092
0.2% Group (n = 15)							
3 min post-challenge	1.16	0.96	1.59	0.99	-0.43	0.75	0.0414
10 min post-challenge	. 1.76	0.86	2.13	0.89	-0.38	0.79	0.0845
20 min post-challenge	1.92	0.86	2.02	1.01	-0.10	1.11	0.7312
0.3% Group (n = 15)							
3 min post-challenge	0.81	0.77	2.00	0.77	-1.19	0.72	<0.0001
10 min post-challenge	1.53	1.14	2.79	0.67	-1.26	1.00	0.0003
20 min post-challenge	1.70	1.11	2.80	0.80	<u>-1.10</u>	1.01	0.0009

nching- 4 Hr Châllenge	LE Eye	LE Eye		Vehicle Eye		Efficacy Score**	
	Mean	Sd.	Mean	Sd.	Mean	Sd.	(2-tail)
0.1% Group (n = 14)							
3 min post-challenge	1.57	1.17	1.82	1.20	-0.25	1.25	0.4883
10 min post-challenge	1.61	0.98	1.93	0.94	-0.32	1.30	0.3700
20 min post-challenge	0.93	1.07	1.36	1.15	-0.43	1.55	0.3212
0.2% Group (n = 16)							
3 min post-challenge	1.69	0.96	1.69	1.08	0.00	0.82	1.0000
10 min post-challenge	1.75	0.93	1.94	1.11	-0.19	0.93	0.4320
20 min post-challenge	0.91	0.78	1.31	1.21	=0.41	0.78	0.0545
0.3% Group (n = 14)							
3 min post-challenge	1.14	0.89	1.86	1.05	-0.71	0.97	0.0168
10 min post-challenge	1.25	0.80	1.89	0.90	-0.64	0.99	0.0302
20 min post-challenge	0.64	1.01	1.21	0.61	-0.57	0.96	0.0438
0.4 411.00	15 500		. dahinin	F.15			
Redness- 4 Hr Challenge	LE Eye	ł	Vehicie	Eye	ETICAC	y Score**	p-vāluē (2-tail)
	Mean	Sđ.	Mean	Sd.	Mean	Sđ.	(Z-tail)
0.1% Group (n - 14)							
3 min post-challenge	0.80	0.77	1.35	0.85	-0.55	0.83	0.0283
10 min post-challenge	1.57	1.02	2.06	0.86	-0.49	0.85	0.0519
20 min post-challenge	1.58	0.87	2.35	0.97	= 0.76	0.97	0.0113
0.2% Group (n - 16)							
3 min post-challenge	0.92	0.91	1.40	0.97	-0.48	0.92	0.0540
10 min post-challenge	1.25	0.87	2.39	0.85	-1.14	0.94	0.0002
20 min post-challenge	1.32	1.02	2.37	1.02	-1.05	1.14	0.0022
0.3% Group (n - 14)							
3 min post-challenge	0.70	0.77	1.25	0.71	-0.55	0.58	8.0038
10 min post-challenge	1.23	1.07	2.31	0.95	-1.08	0.86	0.0004
20 min post-challenge	1.25	1.24	2.27	0.92	-1.02	0.94	0.0013

Visit 3 (Day 21) - Parallel Group	LE Treatment Group N = 16)		Vehicle Treatment Group (N = 15)		Mean Diff. Score	p-value (2-tāii)
	. Mean	Sd.	Mean	Sd.		
Itching						
3 min post-chāllenge	1.55	0.89	2.27	0.96	-0.72	0.0379
10 min post-challenge	1.94	0.90	2.23	0.68	-0.29	0.3144
20 min post-challenge	1.09	0.79	1.50	0.91	=0.41	0.1930
Mean Redness						
3 min post-challenge	1.05	0.89	1.79	0.81	-0.74	0.0145
10 min post-challenge	1.86	0.87	2.19	0.94	-0.33	0.3194
20 min post-challenge	1.88	0.93	2.04	0.90	-0.18	0.6231
Ciliary						
3 min post-challenge	0.89	0.88	1.62	0.82	-0.73	0.0244
10 min post-challenge	1.69	0.80	2.17	1.02	-0.48	0.1547
20 min post-challenge	1.77	0.92	1.98	0.98	÷0.22	0.5293
Conjunctival						
3 min post-challenge	1.17	0.88	1.78	0.86	-0.61	0.0802
10 min post-challenge	1.95	0.91	2.23	0.94	-0.28	0.4076
20 min post-challenge	1.94	0.96	2.10	0.88	-0.16	0.6259
Episcleral						
3 min post-challenge	1.08	0.96	1.70	0.79	-0.62	0.0591
10 min post-challenge	1.94	0.92	2.17	0.90	-0.23	0.4891
20 min post-challenge	1.92	0.95	2.03	0.89	=0.11	0.7386
Chemosis						
3 min post-challenge	0.28	0.34	0.58	0.39	-0.30	0.0278
10 min post-challenge	0.69	0.47	0.85	0.53	-0.16	0.3745
20 min post-challenge	0.73	0.60	0.83	0.64	-0.10	0.6602
Lid Swelling						
3 min post-chállenge	0.20	0.40	0.17	0.36	+0.03	0.7926
10 min post-challenge	0.28	0.45	0.30	0.48	-0.02	0.9114
20 min post-challenge	0.31	0.49	0.28	0.43	+0.03	0.8616
Tearing						
3 min post-challenge	0.00	0.00	0.30	0.80	-0.30	0.1671
10 min post-challenge	0.00	0.00	0.27	0.80	-0.27	0.2170
20 min post-challenge	0.03	0.13	0.38	1.00	-0.35	0.1983

The efficacy of the 0.5% loteprednol appears to be approximately equal to the efficacy observed for the 0.2% and 0.3%.

Visit 4 (Day 35) Parallel Group	LE Treatment Group (N = 16)		Vehicle Treatment Group (N = 14)		Mean Diff, Score	p-value (2-tāil)
	Mean	Sd.	Mean	Sd.		
kching						
3 min post-challenge	1.05	0.95	2.38	0.79	-1.33	0.0003
10 min post-challenge	1.47	0.76	2.05	0.79	-0.58	0.0482
20 min post-challenge	0.59	0.58	1.07	0.90	-0.48	0.0895
Mean Redness				3.30		0.0053
3 min post-challenge	0.67	0.67	1.47	1.01	-0.80	D 0454
10 min post-challenge	1.51	0.84	2.13	0.84	-0.62	0.0151 0.0523
20 min post-challenge	1.62	0.83	2.15	0.99	-0.53	0.0523 0.11 98
Ciliary		-	4	0.00	-0.00	0.1180
3 min post-challenge	0.47	0.69	1.32	1.07	0.05	
10 min post-challenge	1.52	0.81	2.00	0.95	-0.85 -0.48	0.0141
20 min post-challenge	1.59	0.83	2.04	1.14	-0.45	0.1427 0.2308
Conjunctival		0.00	2.04	1.14	-0.45	0.2308
3 min post-challenge	0.80	0.68	1 50	0.03		
10 min post-challenge	1.55	0.82	1.59 2.20	0. 97 0.82	-0.79	0.0143
20 min post-challenge	1.69	0.82	2.25	0.82 0. 9 6	-0.65 -0.56	0.0384
Episcleral		U.UZ	2.27	0.80	-0.50	0.0944
3 min post-challenge	0.76	^				
10 min post-challenge	0.75 1.47	0.71	1.50	1.00	-0.75	0.0235
20 min post-challenge	1.58	0.92 0. 9 0	2.20 2.18	0.79	-0.73	0.0285
Chemosis	1.30	0.90	2.15	0. 9 2	-0.60	0.0819
3 min post-challenge 10 min post-challenge	0.14	0.27	0.38	0.48	-0.24	0.0947
20 min post-challenge	0.55	0.48	0.91	0.50	-0.36	0.0522
·	0.53	0.47	0.82	0.58	-0.29	0.1408
Lid Swelling						
3 min post-challenge	0.06	0.17	0.21	0.47	-0.15	0.2684
10 min post-challenge	0.28	0.41	0.38	0.53	-0.10	0.5868
20 min post-challenge	0.28	0.41	0.36	0.49	-0.08	0.6457
Tearing						
3 min post-challenge	0.00	0.00	0.18	0.67	-0.18	0.3358
10 min post-challenge	0.00	0.00	0.18		-0.18	0.1739
20 min post-challenge	0.00	0.00	0.04	0.13	-0:04	0.3356

There is very little difference between evaluations at Day 21 and Day 35.

ЮР (mmH	g)					LE Eye			Vehicle	e Eye	
Visit				Da	у п		Mean	SD	Range (mm Hg)	Mean	SD	Range (mm Hg)
0.1%	treati	ner	nt group	3								-
2				7	2	7	15.48	2.46	12 - 21	15.59	2.61	12 - 21
2 3				21	2		15.25	2.46	11 - 20	15.18	2.55	11 -20
4 (2 h	r)			35	1:		15.62	2.47	11 - 18	14.31	3.28	8 - 18
4 (4 h	r)			35	14		14.57	1.87	12 - 21	14.43	2.10	10 - 18
0.2%	treatr	nen	nt group)								
2				7	3'	1	15.65	2.25	10 - 21	15.52	2.41	10 - 21
3				21	31		15.87	2.75	10 - 21	15.68	2.60	10 - 20
4 (2 fi	•			35	15	5	15.87	2.17	10 - 22	15.53	2.23	12 - 20
4 (4 hi	7)			35	16	3	16.19	3.53	10 - 20	15.81	2.93	11 - 20
0.3% t	reatn	nen	t group)					•			
2			_ •	7	29)	15.47	2.73	10 - 20	16.14	2.71	10 - 22
3				21	29	}	16.28	2.62	12 - 22	16.10	2.72	12 - 22
4 (2 hr				35	15	5	17.73	3.26	12 - 24	18.40	2.44	12 - 20
4 (4 hr)			35	14	1	16.14	2.66	12 - 20	15.21	2.81	12 - 20
IOP					1 F T		•					
IOF					LE Treat	ıment (Group	•	/ehicle Tre	atment	Group	
			OD		os			OD		OD		
Visit	Day	n	Mean	SD	Mean	SD	Range	n Mean	SD	Mean	SD	Range
2	7	16	15.75	3.11	15.73	3.06	10 - 20	15 16.60	1.45	16.73	1.87	15 - 20
3	21		17.00	2.03	17.00	1.71		15 16.13	2.53	16.67	2.16	11 - 20
4 (2 hr)	35	16	17.50	2.34	17.69	2.27		14 15.67	3.59	16.07	3.83	10 - 22

Intraocular pressures are generally higher in the loteprednol group compared to the vehicle group after 14 days of treatment or more.

Adverse Experiences

	LE 0.1%	.		
D	Eye	Medical Event	Date of Onset	Intensity
5103	άO	COLD	11/05/95	MILD
5123	OD	HEADACHE	11/17/95	MILD
5126	OS	ASTHMA	10/17/96	MILD
5135	OD GO	HEADACHE	10/01/95	MILD
5135	OD	HEADACHE	10/03/95	MILD
5135	OD	HEADACHE	11/04/95	MILD
	os	HEADACHE	11/02/95	MILD
5143	OS			MILD
5143		HEADACHE	11/05/95	
5143	os	COLD	11/07/95	MILD
5143	os	HEADACHE	11/07/95	MILD
5148	os	HEADACHE	11/05/95	MODERATE
5149	0\$	HEADACHE	11/18/95	MODERATE
5150	os	INTERMITTENT ITCHING OU	10/29/95	MILD
5150	OS	INTERMITTENT TEARING OU	10/29/95	MILD
5151	OD	STREP THROAT	11/14/95	MODERATE
5173	os	HEADACHE	10/31/95	MILD
5173	OS	HEADACHE	11/19/95	MODERATE
5176	OD	COLD	11/15/95	MILD
5182	os	HEADACHE	10/30/95	MILD
5190	OD	WHEEZING	11/01/95	MILD
5192	OD	COLD	10/31/95	MILD
5201	os	EYELID SWELLING RAL	11/23/95	MILD
5201	os	DRYNESS R/L	10/23/95	MILD
5201	ÖS	ECZEMA	11/15/95	MILD
5206	QO	JOINT DETERIORATION	10/07/95	MODERATE
		2 DEGREE TRAUMA	1001100	
		E DEGILLE HOLOMA		
	LE 02%			
-		Madical Event	Date of Occasi	Andronalis,
ID .	Eye	Medical Event	Date of Onset	Intensity
5105	Eye OD	TOOTHACHE	10/18/95	MODERATE
5105 5105	Eye OD OD	TOOTHACHE TOOTHACHE	10/18/95 10/02/95	MODERATE MODERATE
5105 5105 5105	Eye OD OD OD	TOOTHACHE TOOTHACHE HEADACHE	10/18/95 10/02/95 11/01/95	MODERATE MODERATE MODERATE
5105 5105 5105 5105	Eye OD OD OD OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95	MODERATE MODERATE MODERATE MODERATE
5105 5105 5105 5105 5111	Eye OD OD OD OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95	MODERATE MODERATE MODERATE MODERATE MILD
5105 5105 5105 5105 5111 5127	Eye OD OD OD OD OD OD OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95	MODERATE MODERATE MODERATE MODERATE MILD MILD
5105 5105 5105 5105 5111 5127 5127	Eye OD OD OD OD OD OD OS OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD
5105 5105 5105 5105 5111 5127 5127 5127	Eye OD OD OD OD OD OD OS OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 10/03/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127	Eye OD OD OD OD OD OD OS OS OS OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL	10/18/95 10/02/95 11/01/95 11/04/95 10/28/95 10/18/95 10/18/95 10/18/95 11/15/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139	Eye OD OD OD OD OD OD OS OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 10/03/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127	Eye OD OD OD OD OD OD OS OS OS OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/28/95 10/18/95 10/18/95 10/18/95 11/15/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139	Eye OD OD OD OD OD OS OS OS OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 10/19/95 11/15/95 10/16/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139	Eye OD OD OD OD OD OS OS OS OS OD OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/19/95 11/15/95 10/16/95 11/16/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153	Eye OD OD OD OD OD OS OS OS OS OD OD OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 10/03/95 11/15/95 11/18/95 11/18/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153	Eye OD OD OD OD OD OS OS OS OS OD OD OD OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 11/15/95 11/18/95 11/18/95 11/11/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5139 5139 5153 5153	Eye OD OD OD OD OS OS OS OS OS OD OD OD OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 10/13/95 11/15/95 11/16/95 11/11/95 11/11/95 10/29/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5153 5164 5164	Eye OD OD OD OD OS OS OS OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/19/95 10/03/95 11/15/95 11/18/95 11/11/95 11/11/95 10/29/95 10/17/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5139 5139 5153 5153 5153 5164	Eye OD OD OD OD OS OS OS OS OD OD OD OD OD OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE HEADACHE HEADACHE COLD	10/18/95 10/02/95 11/01/95 11/04/95 10/28/95 10/18/95 10/18/95 11/15/95 10/16/95 11/11/95 11/11/95 11/11/95 10/17/95 10/17/95 11/04/95 11/04/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5128 5139 5153 5153 5153 5164 5164 5180	Eye OD OD OD OD OS OS OS OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/16/95 11/15/95 10/16/95 11/11/95 11/11/95 11/11/95 10/29/95 10/17/95 11/04/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5153 5153 5153 5164 5186 5186 5185	Eye OD OD OD OD OS OS OS OD OD OD OS OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/18/95 11/15/96 10/16/95 11/11/95 11/11/95 11/11/95 10/29/95 10/13/95 10/13/95 10/13/95 10/20/95 11/14/95	MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5164 5164 5180 5185 5185 5185	Cyn OD OD OD OD OS OS OS OS OD OD OD OS OS OS OD OD OS OS OS OD OD OS OS OD OD OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE COLD	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/18/95 11/15/95 11/11/95 11/11/95 11/11/95 11/11/95 11/11/95 11/11/95 11/11/95 11/13/95 11/14/95 10/13/95 10/16/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5164 5164 5164 5165 5185 5185	Cyn OD OD OD OD OS OS OS OS OD OD OD OS OS OS OD OD OS OS OD OD OS OD OD OS OD OD OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE COLD HEADACHE COLD HEADACHE COLD HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/18/95 11/15/96 10/18/95 11/11/95 11/11/95 11/11/95 11/11/95 11/11/95 11/13/95 10/13/95 11/14/95 10/16/95 10/16/95 10/16/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5128 5139 5153 5153 5153 5164 5164 5180 5185 5185 5185 5185 5186 5187	Eye OD OD OD OD OD OS OS OS OS OS OD OD OD OS OS OS OD OD OS OS OD OD OS OD OS OD OD OS OD OD OS	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE HEADACHE COLD HEADACHE COLD HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/18/95 11/15/95 11/11/95 11/11/95 11/11/95 10/29/95 10/13/95 11/11/95 10/13/95 10/13/95 10/13/95 10/13/95 10/13/95 10/13/95 10/13/95 10/13/95 10/13/95 10/13/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5153 5153 5153 5164 5164 5186 5185 5185 5185 5187 5196 5200 5207	Eye OD OD OD OD OD OS OS OS OS OD OD OD OS OS OS OD OD OS OS OD OD OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/29/95 10/18/95 10/18/95 10/18/95 10/18/95 11/15/95 11/11/95 11/11/95 10/29/95 10/13/95 10/13/95 10/13/95 10/13/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5153 5164 5164 5186 5185 5185 5185 5185 5185 5186 5187 5196 5200 5207	Eye OD OD OD OD OS OS OS OS OS OD OD OD OS OS OS OD OD OD OS OD OD OS OD OD OD OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE HEADACHE COLD HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/18/95 10/18/95 10/18/95 10/18/95 11/15/95 10/18/95 11/11/95 11/11/95 11/11/95 11/11/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5164 5164 5180 5185 5185 5185 5186 5187 5196 5207 5207	Eye OD OD OD OD OD OS OS OS OS OD OD OD OS OS OD OD OD OS OD OD OD OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/28/95 10/18/95 10/18/95 10/18/95 11/15/96 10/16/95 11/11/95 11/11/95 11/11/95 10/18/95 11/11/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 11/18/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5164 5186 5186 5186 5186 5186 5187 5196 5207 5207 5207	Eye OD OD OD OD OD OS OS OS OS OD OD OD OS OS OD OD OD OS OD OD OD OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE COLD HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/28/95 10/18/95 10/18/95 10/18/95 10/18/95 11/15/95 11/11/95 11/11/95 11/11/95 11/13/95 10/13/95 10/13/95 10/13/95 10/18/95 10/18/95 10/18/95 10/18/95 11/14/95 10/18/95 10/18/95 11/14/95 10/18/95 11/16/95 11/16/95 11/16/95 11/17/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5105 5105 5105 5105 5105 5111 5127 5127 5127 5127 5139 5139 5153 5153 5164 5164 5180 5185 5185 5185 5186 5187 5196 5207 5207	Eye OD OD OD OD OD OS OS OS OS OD OD OD OS OS OD OD OD OS OD OD OD OS OD	TOOTHACHE TOOTHACHE HEADACHE HEADACHE MILD COLD SINUS CONGESTION CHEST CONGESTION SINUS CONGESTION CHALAZION RUL BACKACHE HEADACHE ASTHMA SINUS CONGESTION ASTHMA HEADACHE HEADACHE COLD HEADACHE	10/18/95 10/02/95 11/01/95 11/04/95 10/28/95 10/18/95 10/18/95 10/18/95 11/15/96 10/16/95 11/11/95 11/11/95 11/11/95 10/18/95 11/11/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 11/18/95	MODERATE MODERATE MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD

	LE 0.3	3%		
ID	Eye	Medical Event	Date of Onset	Intensity
5117	OD	CONGESTION	11/17/95	MILD
5117	OD	BACKACHE	11/04/95	MILD
5117	OD	BACKACHE	11/08/95	MILD
5119	os	SINUS CONGESTION	10/19/95	MILD
5121	os	SUBDERMAL HEMORRHAGE OD	11/18/95	MILD
5134	OD	HEADACHE	11/03/95	MILD
5145	os	URINARY TRACT INFECTION	11/03/95	MODERATE
5146	os	INDIGESTION	10/17/95	MILD
5146	OS	CAKING OF LASHES	10/23/95	MILD
5146	OS	HEADACHE	11/11/95	MODERATE
5152	OD	HEADACHE	10/17/95	MILD
5152	OD	BACKACHE	11/04/95	MILD
5152	OD	BACKACHE	11/15/95	MILD
<u>5</u> 157	<u> </u>	HEADACHE	11/03/95	MODERATE
5158	ÓŚ	HĚÁĎÁCHĚ	11/18/95	MILD
5157	os	CRAMPS (MENSTRUAL)	10/18/95	MILD
5158	os	HEADACHE	10/31/95	MILD
5158	os	HEADACHE	11/07/95	MILD
5158	os	HEADACHE	11/16/95	MILD
5158	os	HEADACHE	11/18/95	MILD
5172	OD	ASTHMA-WHEEZING	10/23/95	MILD
5189	OD	HEADACHE	10/21/95	MILD
5189	OD	HEADACHE	10/25/95	MODERATE
5189	QO	HEADACHE	10/30/95	MODERATE
5189	OD	HEADACHE	10/22/95	MILD
5189	OD	HEADACHE	11/12/95	MILD
5189	OD	HEADACHE	11/16/95	MILD
5206	OD	SINUS CONGESTION	10/22/95	MILD
5211	OS	COLD	11/17/95	MILD
_	LE 0.5			
ID.	Eye	Medical Event	Date of Onset	Intensity
5106	Eye QD	Medical Event COLD	Date of Onset 10/16/95	Intensity MILD
5106 5106	Eye OD OD	Medical Event COLD BODY ACHES		
5106 5106 5124	Eye OD OD OS	Medical Event COLD BODY ACHES HEADACHE	10/16/95 10/27/95 10/18/95	MILD MILD MILD
5106 5106 5124 5128	Eye OD OD OS OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95	MILD
5106 5106 5124 5128 5131	Eye OD OD OS OS OD	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD	10/16/95 10/27/95 10/16/95 10/30/95 11/16/95	MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142	Eye OD OD OS OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/03/95	MILD MILD MILD MILD MILD MODERATE
5106 5106 5124 5128 5131 5142 5155	Eye OD OD OS OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/03/95	MILD MILD MILD MILD MILD MODERATE MODERATE
5106 5106 5124 5128 5131 5142 5155 5155	Eye OD OD OS OS OD OS OD	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE HEADACHE SINUS CONGESTION	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/03/95 11/16/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155	Eye OD OS OS OD OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/03/95 11/18/95 10/16/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MODERATE
5106 5106 5124 5128 5131 5142 5155 5155 5155	Eye OD OS OS OD OS OD OS OD OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/16/95 10/28/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MODERATE MILD MODERATE MILD
5106 - 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155	Eye OD OS OS OD OS OD OS OD OS OS OS OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/16/95 10/23/95 10/23/95	MILD MILD MILD MILD MILD MILD MODERATE MODERATE MILD MODERATE MILD MODERATE MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155 5155	Eye OD OD OS OS OD OS OD OS OS OS OS OS OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/28/95 10/23/95 11/02/95	MILD MILD MILD MILD MILD MILD MODERATE MODERATE MILD MODERATE MILD MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155 5156 5159 5160	Eye OD OD OS OS OD OS OD OS OS OD OS OS OD OS OS OS OS OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/16/95 10/23/95 11/102/95 10/23/95	MILD MILD MILD MILD MILD MILD MODERATE MODERATE MILD MODERATE MILD MILD MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155 5156 5156 5160 5160	Eye OD OD OS OS OD OS OD OS OS OD OS OS OS OS OS OS OS OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/16/95 10/23/95 10/23/95 10/27/95 11/02/95	MILD MILD MILD MILD MILD MILD MODERATE MODERATE MILD MODERATE MILD MILD MILD MILD MILD MILD MILD
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5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5156 5160 5160 5160	Eye OD OD OS OS OD OS OS OS OS OS OS OS OS OD OS OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE HEADACHE HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/16/95 10/23/95 10/23/95 11/02/95 11/02/95 11/02/95 11/02/95 11/02/95 11/02/95 11/02/95 11/02/95 11/02/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
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5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5156 5160 5160 5160	Eye OD OD OS OS OD OS OS OS OS OS OS OS OS OS OD OS OS OD OS OS OD OD OD	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE	10/16/95 10/27/95 10/18/95 10/18/95 11/18/95 11/18/95 11/18/95 10/18/95 10/23/95 10/23/95 11/02/95 11/02/95 11/03/95 11/03/95 10/23/95 11/03/95 10/23/95 10/23/95 10/23/95 10/23/95 10/23/95 10/18/95 10/18/95 10/18/95 10/18/95	MILD MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5156 5160 5160 5160	Eye OD OD OS OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE HEADACHE COLD HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE	10/16/95 10/27/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/28/95 10/23/95 11/02/95 11/02/95 11/02/95 11/08/95 10/23/95 11/08/95 10/23/95 10/23/95 10/23/95 10/23/95 10/18/95 10/18/95 10/18/95 10/19/95 11/13/95 10/15/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5156 5160 5160 5160	Eye OD OD OS OS OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE HEADACHE	10/16/95 10/27/95 10/18/95 10/18/95 11/18/95 11/18/95 11/18/95 11/18/95 10/28/95 10/28/95 11/028/95 11/028/95 11/08/95 11/08/95 11/08/95 11/08/95 10/28/95 10/28/95 10/28/95 10/28/95 10/28/95 10/28/95 10/28/95 10/28/95 10/28/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
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5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155 5156 5160 5160	Eye OD OD OS OS OD OS OS OS OS OS OS OS OS OS OD OD OD OD OD OD OD	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE COLD HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE	10/16/95 10/27/95 10/18/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/26/95 10/23/95 10/23/95 11/02/95 11/03/95 10/23/95 11/03/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155 5156 5160 5160	Eye OD OD OS	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE	10/16/95 10/27/95 10/18/95 10/18/95 10/18/95 11/18/95 11/18/95 11/18/95 10/18/95 10/28/95 10/23/95 11/02/95 11/02/95 11/03/95 10/19/95 11/03/95 10/19/95 11/02/95 10/19/95 11/02/95 11/02/95 10/19/95 11/02/95 11/02/95 11/02/95 10/15/95 10/15/95 10/15/95 11/03/95 10/15/95 10/15/95 10/15/95 10/15/95 10/15/95 10/15/95 10/15/95 10/15/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD
5106 5106 5124 5128 5131 5142 5155 5155 5155 5155 5155 5156 5160 5160	Eye OD OD OS OD	Medical Event COLD BODY ACHES HEADACHE HEADACHE COLD HEADACHE SINUS CONGESTION ECZEMATOUS RASH SINUS CONGESTION HEADACHE KNEE PAIN HEADACHE HEADACHE COLD HEADACHE COLD HEADACHE CRUSTING ON EYELIDS IN AM HEADACHE SINUS CONGESTION	10/16/95 10/27/95 10/18/95 10/18/95 10/30/95 11/18/95 11/18/95 11/18/95 10/26/95 10/23/95 10/23/95 11/02/95 11/03/95 10/23/95 11/03/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 10/18/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95 11/13/95	MILD MILD MILD MILD MILD MODERATE MODERATE MILD MILD MILD MILD MILD MILD MILD MILD

5174	OD	HEADACHE	11/10/95	MILD
5183	OS	HEADACHE	10/24/95	MILD
5183	os	OCULAR IRRITATION	11/04/95	MILD
5183	OS	BLURRED VA	11/04/95	MILD
5188	OD	SINUS CONGESTION	10/15/95	MILD
5188	OD	HEADACHE	10/21/95	MILD
5188	OD	HEADACHE	11/01/95	MILD
5186	OD	HEADACHE - SINUS	11/13/95	MODERATE
5199	OS	HEADACHE	11/01/95	MODERATE
5209	OD	ALLERGIC CONJUNCTIVITIS	11/05/95	MILD
5216	os	HEADACHE	11/18/95	MODERATE
5217	os	BACK PAIN	10/18/95	MODERATE
5217	os	BACK PAIN	10/30/95	MILD
5217	O\$	HEADACHE	11/07/95	MODERATE
5217	OS	HEADACHE	11.09/95	MILD
5217	os	HEADACHE	11/15/95	MILD

Due to the design of the study, few adverse events are expected. No new events were observed in this study.

APPEARS THIS WAY ON ORIGINAL

9. Reviewer's Overview of Efficacy

Marginal efficacy has been demonstrated in the resolution of itching and redness. This constitutes a marginal demonstration of the efficacy for the relief of signs and symptoms of seasonal allergic conjunctivitis. Taken with the studies submitted for NDA 20-583, adequate efficacy has been demonstrated.

10. Reviewer's Overview of Safety

The total number of patients studied with the loteprednol etabonate ophthalmic suspension, 0.2% is too small by itself to establish safety, however, taken with the patients studied with the 0.3%-0.5%, adequate safety has been established for use in the relief of signs and symptoms of allergic conjunctivitis.

Adverse experiences in the limited studies (duration and number of patients) were generally confined to mild to moderate ocular events. There was an increased chance of increased IOP during use.

APPEARS THIS WAY ON ORIGINAL

THIS SECTION WAS DETERMINED NOT TO BE RELEASABLE



12 Conclusions

The submitted studies in NDA 20-803 taken together with the studies in NDA 20-583 and NDA 20-841 demonstrate safety and efficacy for the temporary relief of the signs and symptoms of seasonal allergic conjunctivitis.

13 Recommendations

- 1. Following resolution of any chemistry/manufacturing issues and labeling issues, NDA 20-583 is recommended for approval for the temporary relief of the signs and symptoms of seasonal allergic conjunctivitis. Approval for the steroid class indication is not recommended.
- 2. The applicant should submit revised labeling consistent with the recommendations in this review.
- 3. The proposed tradename should be specified.
- 4. Table 4 in Clinical Study Report 145 (Volume 17, Page 55) should be corrected. The standard deviations are in error.
- 5. The pH range and other specifications in the NDA summary differs from other sections of the NDA [Table 2.5.2.3 vs Table 2.5.2.5]. The specifications should be clarified and be consistent.
- 6. Issues related to water loss and the formation of "aggregate" material after storage of inverted containers will need to be resolved prior to approval.
 - A. What is the aggregate composed of?
 - B. Does the aggregate recombine with the suspension on shaking? If so, how quickly?
 - C. Can the aggregate be cleared by dispensing a couple of drops of the suspension? If so, does this affect the composition of the rest of the suspension?

Wiley A. Chambers, M.D. Medical Officer, Ophthalmology

ce: NDA 20-803
HFD-550
HFD-340
HFD-550/PM/LoBianco
HFD-830/CHEM/Feñselaü
HFD-805/MICRO/Cooney
HFD-550/PHARM/Weir
HFD-550/MO/Chambers